FORM PTO-1390 (REV 10-2000) U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMA'K OFFICE

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TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371

B-4075PCT 618484-4

U.S. APPLICATION NO. (If Known, see 17 CFR 1.5)
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INTERNATIONAL APPLICATION NO. PCT/IT99/00226

INTERNATIONAL FILING DATE 19 July 1999 PRIORITY DATE CLAIMED 17 July 1998

TITLE OF INVENTION A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS, GELLS

AND TRANSGENIC PLANTS CONTAINING IT	
APPLICANT(S) FOR DO/EO/US Corrado FOGHER	
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:	
1. This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.	
2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371.	
3. X This is an express request to promptly begin national examination procedures (35 U.S.C. 371(f)).	
4. X The US has been elected by the expiration of 19 months from the priority date (PCT Article 31).	
5. X A copy of the International Application as , published.	
a. is attached hereto (required only if not communicated by the International Bureau).	
b. As been communicated by the International Bureau.	
c. is not required, as the application was filed in the United States Receiving Office (RO/US).	
6. An English language translation of the International Application as filed (35 U.S.C. 371(c)).	
7. Amendments to the claims of the International Application under PCT Article 19(35 U.S.C. 371(c)(3))	
a. are attached hereto (required only if not communicated by the International Bureau).	
b. Have been communicated by the International Bureau.	
c. L have not been made; however, the time limit for making such amendments has NOT expired.	
d. X have not been made and will not be made.	
8. An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C.371(c)(3))	).
9. An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).	
An English language translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).	
Items 11 to 16 below concern document(s) or information included:	
11. An Information Disclosure Statement under 37 CFR 1.97 and 1.98.	
12. An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is	included.
3. 🔀 A FIRST preliminary amendment, with copy of amended claims.	
A SECOND or SUBSEQUENT preliminary amendment.	
4. A substitute specification.	
15. A change of power of attorney and/or address letter.	
6. XX Other items or information:	
Copy of Form PCT/IB/308 Copy of PCT Request Copy of PCT Demand Copy of Form PCT/IPEA/416 with International Preliminary Examination Report with annexes Copy of International Search Report (see Information Disclosure Statement)	
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17. The following fees are submitted:				CA	LCULATION		PTO USE ONLY	
BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5)):								
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Applicant claims small entity status. See 37 CFR 1.27. The fees indicated above are reduced by 1/2.						772.00	4	
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TOTAL NATIONAL FEE =					\$	772.00		
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property					\$			
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a. A check in the amount of \$ 772.00 to cover the above fees is enclosed.								
b. Please charge my Deposit Account No in the amount of \$ to cover the above fees.  A duplicate copy of this sheet is enclosed.						the above fees.		
c. The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 12-0415. A duplicate copy of this sheet is enclosed.								
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Telephone No.: (323) 934-2300			RATION NUMBER					
тетегах	no.: (323) 934	0202		KEGISTR	A FION	NUMBER		

Applicant or Patentee <u>Corrado FOGHER</u> <u>* Attorney's Docket No. <u>B-4075PCT 618484-4</u> Serial or Patent No. <u>09/743,823</u></u>
Filed or Issued: 16 January 2001
International Application No. PCT/IT99/00226 filed 19 July 1999
For: "A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS,
CELLS AND TRANSGENIC PLANTS CONTAINING IT"
Verified Statement (Declaration) Claiming Small Entity Status
37 CFR 1.9(f) and 1.27(c) - Small Business Concern
the owner of the small business concern identified below:
an official of the small business concern empowered to act on behalf of the concern identified below:
Name of Concern: PLANTECHNO S.r.I.  Address of Concern: Via Staffolo 60 26040 Casalmaggiore CR ITALY
I hereby declare that the above identified small business concern qualifies as a small business concern as defined in 13 CFR 121.3-18, and reproduced in 37 CFR 1.9(d), for purposes of paying reduced fees under Section 41(a) and (b) of Title 35, United States Code, in that the number of employees of the concern, including those of its affiliates, does not exceed 500 persons. For purposes of this statement, (1) the number of employees of the business concern is the average over the previous fiscal year of the concern of the persons employed on a full-time, part-time or temporary basis during each of the pay periods of the fiscal year, and (2) concerns are affiliates of each other when either, directly or indirectly, one concern controls or has the power to control the other, or a third party or parties controls or has the power to control both.
I hereby declare that rights under contract or law have been conveyed to and remain with the small business concern identified above with regard to the Invention, entitled: "A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT" by inventor(s): Corrado FOGHER
described in:
[ ] the specification filed herewith. [X] application serial no.: 09/7¥3823, filed 16 January 2001 [ ] patent no, issued
If the rights held by the above identified small business concern are not exclusive, each individual, concern or organization having rights to the invention is listed below*, and no rights to the invention are held by any person, other than the inventor, who could not qualify as a small business concern under 37 CFR 1.9(c) or by any concern which would not qualify as a small business concern under 37 CFR 1.9(d) or a non-profit organization under 37 CFR 1.9(e). *NOTE: Separate verified statements are required from each named person, concern or organization having rights to the invention averring to their status as small entities. (37 CFR 1.27)
NAME:
ADDRESS
[ ]INDIVIDUAL [ ]SMALL BUSINESS CONCERN [ ] NONPROFIT ORGANIZATION
NAME:
ADDRESS
[ ]INDIVIDUAL [ ] SMALL BUSINESS CONCERN [ ] NONPROFIT ORGANIZATION
I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. [37 CFR 1.28(b)]
I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.
Name of Person Signing: NICOLA FOGHEAL Title of Person Other Than Owner: GENERAL RANAGER
Address of Person Signing: Via TiciNo 32 26041 (ASALTAGGIO NE CR. ITALY
Signature: Date: 2 MORCH 2001
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Applicant or Patentee: Corrado FOGHER Attornéy's Docket No. B-4075PCT 618484-4  Serial or Patent No. 09/743,823  Filed or Issued: 16 January 2001  International Application No. PCT/IT99/00226, filed 19 July 1999  For: "A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT"
VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 CFR 1.9 (f) and 1.27(b)) INDEPENDENT INVENTOR(S)
As a below named inventor, I hereby declare that I qualify as an independent inventor as defined in 37 CFR 1.9 (c) for purposes of paying reduced fees under section 41 (a) and (b) of Title 35, United States Code, to the Patent and Trademark Office with regard to the invention entitled "A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT "described in
[ ] the specification filed herewith. [ ] application serial no.:
i have not assigned, granted, conveyed or licensed and am under no obligation under contact or law to assign, grant, convey or license, any rights in the invention to any person who could not be classified as an independent inventor under 37 CFR 1.9 (c) if that person had made the invention, or to any concern which would not qualify as a small business concern under 37 CFR 1.9 (d) or a nonprofit organization under 37 CFR 1.9 (e).
Each person, concern or organization to which I have assigned, granted, conveyed or licensed or am under an obligation under contract or law to assign, grant, convey or license any rights in the invention is listed below
[ ] no such person, concern, or organization [X] persons, concerns or organizations listed below*  *NOTE: Separate venified statements are required from each named person, concern or organization having nghts to the invention averting to their status as small entities. (37 CFR 1 27)
FULL NAME PLANTECHNO S r L
ADDRESS <u>Via Staffolo 60 26040 Casalmaggiore CR ITALY</u> [] INDIVIDUAL [X] SMALL BUSINESS CONCERN [ ] NONPROFIT ORGANIZATION
FULL NAME
ADDRESS
I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate (37 CFR 1 28 (b))
i hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.
9 7 1
Corrado FOGHER  Nome of First Inventor  Signature of First Inventor  Date

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

) Re: Preliminary Amendment Applicant: Corrado FOGHER ) Group: not yet assigned U.S. Appln. No.: not yet assigned U.S. Filing Date: concurrently ) Examiner: not yet assigned herewith International Application No: PCT/IT99/00226 International Filing Date: Our Ref.: B-4075PCT 618484-4 19 July 1999 For: "A SYNTHETIC POLYNUCLEOTIDE) CODING FOR HUMAN LACTOFERRIN,

VECTORS, CELLS AND TRANSGENIC

PLANTS CONTAINING IT"

) Date: January 16, 2001

Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231

Attn: United States Designated/Elected Office (DO/EO/US)

Sir:

Prior to examination of the above-identified application, it is respectfully requested that the following amendments be made to the Claims as amended during IPE:

### IN THE CLAIMS

- 1. (amended) A polynucleotide coding for human lactoferrin, [characterized in that it has] having a sequence totally or partially corresponding to the sequence reported in the annexed sequence listing as SEQ ID NO:1, said sequence being optimized for the in plant expression.
- 2. (unchanged) The polynucleotide according to claim 1, wherein said polynucleotide has fused at the 5'-terminus the leader sequence of 7s basic globulin gene or the leader sequence of ßconglycinine gene.

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- 3. (unchanged) The polynucleotide according to claim 2, wherein said leader sequence is the sequence reported in the annexed sequence listing as SEQ ID NO: 13 or the sequence reported in the annexed sequence listing as SEQ ID NO: 14.
- 4. (amended) A recombinant DNA vector comprising the polynucleotide according to [any of] claim[s] 1 [to 3] operatively linked to regulation elements allowing the expression of said polynucleotide.
- 5. (amended) The recombinant DNA vector according to claim 4, wherein said regulation elements [is] are a plant expression cassette allowing the tissue specific expression of said polynucleotide.
- 6. (unchanged) The vector according to claim 5, wherein said plant expression cassette includes the promoter of the gene coding for the protein basic globulin 7 S.
- 7. (unchanged) The vector according to claim 6, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:21
- 8. (amended) The vector according to claim 6 [or 7], wherein said plant expression cassette includes the leader sequence of the gene coding for the protein basic globulin 7 S.
- 9. (unchanged) The vector according to claim 8, wherein said leader sequence is the sequence reported as SEQ ID NO: 13.
- 10. (unchanged) The vector according to claim 5, wherein said plant expression cassette includes the promoter of the gene coding

for ß-conglycinine protein.

- 11. (unchanged) The vector according to claim 10, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:22.
- 12. (amended) The vector according to claim 10 [or 11], wherein said plant expression cassette includes the leader sequence of the gene coding for the ß-conglycinine protein.
- 13. (unchanged) The vector according to claim 12, wherein said leader sequence is the sequence reported in the annexed sequence listing as SEQ ID NO: 14.
- 14. (amended) A vegetal cell including the polynucleotide according to [any of] claim[s] 1 [to 3].
- 15. (amended) A vegetal cell including the vector according to [any one of the] claim[s] 4 [to 13].
- 16. (amended) A cellular aggregation obtainable from cells according to claim 14 [or 15].
- 17. (unchanged) The cellular aggregation according to claim 16, wherein said aggregations are calluses capable of regenerating transgenic plants.
- 18. (amended) A transgenic plant including in a tissue cell the polynucleotide according to [any of] claim[s] 1 [to 3].
- 19. (unchanged) The transgenic plant according to claim 18, wherein said tissue cell is a storage tissue cell.

- 20. (unchanged) The transgenic plant according to claim 18, wherein said tissue cell is a fruit tissue cell.
- 21. (amended) The transgenic plant according to [any of] claim[s] 18 [to 20], said plants being selected from the group consisting of solanaceae, cereals, leguminosae, fruit bearing plants and horticultural plants.
- 22. (unchanged) The transgenic plant according to claim 21, said plant being selected from the group consisting of soya, tobacco and rice.
- 23. (amended) [Use of] A method of using the vector according to [any of] claim[s] 4 [to 13] for the transformation of vegetal cells.
- 24. (amended) [Use of] A method of using the transgenic plant according to [any one of the] claim[s] 18 [to 22,] for the production of nutriceuticals.
- 25. (amended) [Use of] A method of using the transgenic plant according to [any of] claim[s] 18 [to 22,] for the production of human lactoferrin.
- 26. (amended) [Use of] A method of using the transgenic plant according to [any one of] claim 18 [to 22,] for the production of lactoferrin flours or of lactoferrin extracts obtained from tissues of said transgenic plant.
- 27. (amended) [Use of] A method of using the transgenic plant according to [any one of the] claim[s] 18 [to 22] for the production of functional foods containing lactoferrin.

- 28. (amended) The [use]method according to claim 27, wherein said functional food being selected from the group consisting of vegetal milks, fruit juices, fruit and/or vegetable homogeneized foods.
- 29. (unchanged) A plant expression cassette allowing the tissue specific expression of a gene of interest comprising the promoter of the gene coding for the protein basic globulin 7 S.
- 30. (unchanged) The plant expression cassette according to claim 29, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:21.
- 31. (amended) The plant expression cassette according to claim 29 [or 30], wherein said plant expression cassette includes the leader sequence of the gene coding for the protein basic globulin 7 S.
- 32. (unchanged) The plant expression cassette according to claim 31, wherein said leader sequence is the sequence reported as SEQ ID NO: 13.
- 33. (amended) A recombinant DNA vector comprising a gene of interest under the control of the plant expression cassette according to [any of] claim[s] 29 [to 32].
- 34. (amended) The vector according to claim 33 [when depending on claim 31 or 32,]wherein said plant expression cassette includes the leader sequence of the gene coding for the protein basic globulin 7 S and wherein said gene of interest is fused to the leader sequence.

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35. (unchanged) A plant expression cassette allowing the tissue specific expression of a gene of interest comprising the promoter of the gene coding for the ß-conglycinine protein.

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- 36. (unchanged) The plant expression cassette according to claim 35, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:22.
- 37. (amended) The plant expression cassette according to claim 35 [or 36], wherein said plant expression cassette includes the leader sequence of the gene coding for the [leader sequence of the gene coding for the] ß-conglycinine protein.
- 38. (unchanged) The plant expression cassette according to claim 37, wherein said leader sequence is the sequence reported as SEQ ID NO: 14.
- 39. (amended) A recombinant DNA vector comprising a gene of interest under the control of the plant expression cassette according to [any of] claim[s] 35 [to 38].
- 40. (amended) The vector according to claim 39 [when depending on claim 36 or 37,] wherein said plant expression cassette includes the leader sequence of the gene coding for the ß-conglycinine protein and wherein said gene of interest is fused to the leader sequence.
- 41. (amended) A vegetal cell including the vector according to [any one of the] claim[s] 33[, 34, 39 or 40].
  - 42. (unchanged) A cellular aggregation obtainable from the

cell according to claim 41.

- 43. (unchanged) The cellular aggregation according to claim 42, said aggregations being calluses capable of regenerating transgenic plants.
- 44. (amended) A transgenic plant including in a tissue cell the vector according to [any of] claim[s] 33[, 34, 39 or 40].
- 45. (unchanged) The transgenic plant according to claim 44, wherein said tissue cell is a storage tissue cell.
- 46. (amended) The transgenic plant according to claim 44, wherein said tissue cell[s] is a fruit tissue cell.
- 47. (amended) The transgenic plant according to [any of] claim [43 to 45]44, said plant[s] being selected from the group consisting of solanaceae, cereals, leguminosae, fruit bearing plants and horticultural plants.
- 48. (unchanged) The transgenic plant according to claim 47, said plant being selected from the group consisting of soya, tobacco and rice.
- 49. (amended) [Use of] A method of using the vector[s] according to claim[s] 33[, 34, 39 or 40] for the transformation of vegetal cells.

Please add the following new claims:

50. A vegetal cell including the vector according to claim 39.

- 51. A cellular aggregation obtainable from the cell according to claim 50.
- 52. The cellular aggregation according to claim 51, said aggregations being calluses capable of regenerating transgenic plants.
- 53. A transgenic plant including in a tissue cell the vector according to claim 39.
- 54. The transgenic plant according to claim 53, wherein said tissue cell is a storage tissue cell.
- 55. The transgenic plant according to claim 53, wherein said tissue cell is a fruit tissue cell.
- 56. The transgenic plant according to claim 53, said plant being selected from the group consisting of solanaceae, cereals, leguminosae, fruit bearing plants and horticultural plants.
- 57. The transgenic plant according to claim 56, said plant being selected from the group consisting of soya, tobacco and rice.
- 58. A method of using the vector according to claim 39 for the transformation of vegetal cells.

### **REMARKS**

The claims in this application were amended during International Preliminary Examination (IPE). A copy of the amended and claims

attached. This Amendment is directed to the claims as amended during IPE and not to the claims as filed. Thus the amendment made during IPE must be entered before this amendment is entered. This Amendment amends Claims 4, 8, 12, 14-16, 18, 21, 23-27, 31, 33-34, 37, 39-41, 44, 47, and 49 so that they are no longer multiply dependent. Claims 50-58 have been added to provide similar coverage. The Applicant may elect to amend Claims 4, 8, 12, 14-16, 18, 21, 23-27, 31, 33-34, 37, 39-41, 44, 47, and 49 to make them again multiply dependent or to add additional claims to this application to provide coverage similar to, broader than or narrower than the present claims once examination on the merits has begun.

This Amendment also amends claims 23-28 and 49 to eliminate non-statutory "use" claims.

The Examiner should ensure that the amendments made by the IPEA are entered together with the amendments set forth above.

Respectfully submitted,

Richard P. Berg Reg. No. 28,145

Attorney for Applicant

LADAS & PARRY

5670 Wilshire Boulevard #2100 Los Angeles, California 90036

(323) 934-2300

Enclosed: copy of Claims as amended during IPE (5 pages)

PCT/IT99/00226 -

JC07 Rec'd PCT/PTO 1 6-JAN 2001

A SYNTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, VECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT

## DESCRIPTION

## Field of the invention

The present invention relates to the field of the vegetable biotechnologies, and in particular to the plants and vegetal cells transformation techniques and systems, to cells and transgenic plants thus generated, and to their use.

## State of the art

In recent years, due to the large variety of applications deriving from the utilization of the genetic engineering techniques in the biology of the vegetals, the use of genetically modified plants has gradually increased.

In particular, the development of techniques for the transformation of plants into organisms capable of producing proteins of commercial interest has acquired a remarkable practical importance.

In fact, the generation of recombinant plants containing a heterologous gene of interest, and their use in production processes on an industrial scale, allows to overcome a series of drawbacks characterizing the production systems presently in use, particularly those based on cell cultures.

Indeed, recombinant DNA technology allowed the generation of transgenic cells that are used in the In production of heterologous proteins of interest. (and specifically mammal) animal particular, cultures allow the production of proteins of interest, even extremely complex ones, in their native form, but expensive, extremely related process is hectoliters of cells are required for the production of an amount of proteins sufficient for a commercial (e.g., pharmacological or alimentary) use (Stowell et al., 1991).

An alternative in this direction is then provided by

the production, carried out in prokaryotic cell cultures, a cheaper process, meeting however a serious obstacle in the inability of those systems to effect the post transcriptional and post-translational modifications required for the expression of complex heterologous proteins, carried out only by the eukaryotic systems (Glick and Pasternak, 1994).

Therefore, a possible solution was pursued in the transformation of complex eukaryotic systems which could ensure at the same time the production of active proteins and the utilization of inexpensive processes. (Watson et al. 1992).

In particular, plants possess the required potential, and their capacity of functioning the production of complex proteins bioreactors for (plants are highly efficient in this type of processes, high degree of expression), ensuring a in a costeffective way (plant cultivation is inexpensive) and with a high yield (each single harvest can yield high amounts of proteins) was proved (Fraley and Schell, 1991).

Further, very many plants satisfy the important requirement of the non-allergenicity needed for production systems of recombinant proteins of pharmacological or alimentary interest. In fact, organisms used in those productions must belong to the GRAS (Generally Recognized As Safe) organism category, i.e. organisms that having been used by man for a long time, are considered safe for man and for the ecosystem as well. Obviously, plants such as Leguminosae, cereals, tobacco, horticultural plants in general and fruit trees, satisfy this requirement in nature. Among leguminosae, Soya is a basic food in the diet of many populations mostly of the Third World Countries, but recently in European Countries as well. In fact, Soya-derived compounds constitute ingredients that are present in a vast number of food products, such as e.g., lecithin and

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seed oil, while the seeds of this Leguminosa yield a flour that is employed in various food like soybean milk containing approximately the 8% thereof.

Therefore plants, while being the raw material from which the product can be extracted according to conventional processes, at the same time constitute an alternative to the conventional production. In fact, they can be used as functional foods, i.e. foods genetically modified so as to be enriched from a nutritional point of view, and in case assuming important properties as a natural drug.

Thus the step of the protein extraction from the plant has been eliminated, and the heterologous protein expression does not prelude to an extraction and purification thereof, steps that account for most of the production cost of a drug, but simply enriches a vegetable nutriment, which thus becomes a nutriceutical, i.e. a nutriment having a pharmaceutical value.

This is the reason for the research efforts aimed at generating genetically modified plants, optimized for the above-mentioned applications.

However, to date the evaluation of the heterologous expression capacity typical of various plant species was exclusively performed under laboratory conditions, or anyhow on relatively limited surfaces in hothouse.

For instance, expression in plants of enkephalin and human serum albumin, as well as of mice monoclonal antibodies was studied (Watson et al., 1992). More recently, always referring to proteins relevant from a pharmacological point of view, two additional human proteins of therapeutic importance, i.e. interleukin-6 and C protein (a serum anticoagulant), were successfully expressed in tobacco. In all these cases the model plant on which the functionality of the prepared gene was tested is tobacco, whereas usually the plant finally selected for production is a leguminosa, whose seed content in storage proteins is high.

A DOMESTIC SECTION AND ADDRESS OF THE SECTION ADD

These experiments proved that often the expression levels of the heterologous protein in the vegetal bioreactors do not reach high enough levels to meet the commercial demands, and that anyhow it can be improved applying new information on the plant gene control. Specifically, it was demonstrated that the level of produced recombinant protein has to reach more than 1% of the total protein amount in order to become economically significant, a level obviously not achieved by the simple introduction of the heterologous gene, whose expression therefore needs to be maximized.

In relation to the vegetal cell biology, in order to maximize the level of the in plant expression, an action on various levels is necessary: increase of genic transcription, increase of transcript stability and of the translation process yield. Moreover, it is further necessary to fix the inserted gene and to minimize the risk of the occurrence of silencing or of genic inactivation. All these factors are crucial in assessing the in plant expression level of the heterologous gene, that, as afore stated, apart from some exceptions usually settles at rather low levels (Owen and Pen, 1996).

Among all these, the most crucial factor, together depending with transcription level preferential presence of certain codons, would seem to be stability of the recombinant protein heterologous host, as the likely probable cause of its easy elimination. The relative instability might be the one hand of the inability of consequence, on translated heterologous protein to assume a on the other hand of structural conformation, ultrastructural compartment in which it is directed after the translation, where the presence of proteases and of particular pH values determine its degradation.

Accordingly, it is therefore important both to provide the modification of the heterologous sequences in order to ensure the codon optimization and to carry out a

 careful selection of the targeting sequences, capable of directing the translated protein into preferential ultrastructural compartments, as i.e. the seed storage tissue, capable of ensuring the stability of the product. Concerning the latter issue, the best strategy of action in this sense, is that of providing the maintenance of the endogenous signal sequences of the plant selected for the final production. In fact, the adoption of these sequences prevents the alteration of the cell internal consequence of the unavoidable accumulation of exogenous proteins that would take place in their absence. In this regard, the fact that the ultrastructural different compartments have characteristics in the cells of the various tissues has to be taken into account.

For instance, the heterologous proteins accumulated together with the storage proteins of the seeds in the transgenic plants, are more stable as compared with those of the vegetative organs (Owen and Pen, 1996). Probably, the reason of such different stability can be found in the different protease activities recorded in the vacuoles of the leaf meristem cells, with respect to those observed in the vacuoles of the seed. It is therefore understandable how research efforts are aimed at the generation of transgenic plants in which the heterologous protein be preferably expressed in the seed (Owen and Pen, 1996).

Furthermore, overall the seed constitutes the vegetal organ most used by man for its ease of conservation and, obviously, for its caloric and protein contribution. The seed consists of the plant embryo enveloped by storage tissues that provide energy and nitrogen during germination. Usually the storage tissue function is mainly effected by the endosperm, but in Leguminosae, as e.g. Soya, the cotyledons develop remarkably, and acquire this function.

The storage function for the nitrogen component is

carried out by particular proteins accumulated in the protein bodies, inside compartments in the endocellular membrane systems. The amount of proteins in the seed varies from about 10-15% in cereals to about 25-35% in Leguminosae, therefore seeds are an important protein source in the human diet (Shewry et al., 1995).

In order to exploit the expression system of the seed storage proteins, first of all it has to considered that the storage proteins of all plants have some functional and physiological common characteristics: their synthesis is controlled during the seed ripening according to the nutritional needs, and they are stored in protein bodies. In particular, the Leguminosae storage proteins are divided in two classes: globulins and lectins. Globulins are the most widespread storage proteins, present not only in leguminosa but monocotyledons and in other dicotyledons as well. globulin class in turn is subdivided into two subclasses: legumins (11S hexameric proteins) and vicilins trimeric proteins). Also  $\beta$ -conglycinine and basic globulin, whose regulation elements were used to perform tissue-specific expression in seed (Shewry et al., 1995) belong to the vicilin subclass, but whereas the former studied in detail, no detailed information available on the basic globulin functioning.

 $\beta$ -conglycinine is a storage protein of the Soya seed (Glycine max.), consisting of three different subunits,  $\alpha$ ,  $\alpha$ ',  $\beta$  that interact non-covalently to form a trimer complex. The subunits are coded by a multigene family of 15 genes grouped in six nuclear DNA regions, whose expression is strictly regulated so as to be modulated during the plant life (Harada et at. 1989). Control is tissue-specific as well as stage-specific. In fact, the expression of each subunit is activated at high levels at the moment of embryo development, from the heart shaped phase until complete ripening, whereas it is repressed before the dormancy phase.

Moreover, expression occurs exclusively on specific plant zones, like cotyledons, according to an expression pattern that varies in the course of their ripening; at first it is activated on the outer cotyledon area, then a wave distribution from the outside to the inside is observed and lastly it turns out to be uniformly spread over the entire cotyledon tissue. However, during the heart shaped phase (18 days after pollination), the gene encoding the  $\beta$ -conglycinine also expresses itself in the embryonal axis, whereas it does not express in endosperm, tegmen and in already differentiated tissues (Perez-Grau and Goldberg, 1989). The same behavior also occurs in the seed of transgenic plants belonging to other families, as e.g., tobacco, proving the same control mechanism to be functional in solanaceae as well (Naito et al. Perez-Grau and Goldberg, 1989).

regulation of subunits  $\alpha/\alpha$ ß and genic expression occur at a transcriptional as well as a posttranscriptional level (Harada et al., 1989).  $\alpha$  subunit, of 76 Kd, with a 2.5 Kb mRNA, is accumulated more precociously and in a larger amount as compared to  $\boldsymbol{\beta}$ subunit. This behavior is due to the greater strength of the  $\alpha$ ' subunit due to the presence of an enhancer region, in  $\beta$ , and of a sequence stabilizing the  $\alpha'$ transcript, it also absent in  $\beta$  (probably a 560 pb region in the first exon of the  $\alpha'$  transcript) (Harada et al., 1989). This difference in expression has also been highlighted in transformed tobacco plants (Bray et al., 1987).

In contrast with the  $\alpha'$  subunit, the expression level of the  $\beta$  subunit, is also influenced by abiotic stresses, methionine level in the ground and presence of ABA. The base elements involved in gene control at transcriptional level for the  $\alpha'$  subunit are clustered in the 905 pb region at 5' of the transcription start site, region called URS (Upstream Regulatory Sequence). Inside this area specific sequences functioning as site-specific

enhancers have been detected. Among these, the legumin boxes (5'-CATGCAC.3' and 5'-CATGCAT-3'), elements that are found in many other genes encoding seed-specific proteins, in particular in legumes. A coordinated action of the two sequences determines a 10-fold increase of the seed gene expression level. The regulation by the above elements seems to be of a positive kind (Chamberland et al. 1992), however so far trans-elements specifically interacting in those sites have not yet been found. Sitespecific expression also requires the coordinated action of elements operating in cis, not yet characterized, located in the region at 5' of the legumin boxes and at 3' of the promoter (Chamberland et al., 1992). Probably the region at 5' includes a negative control site, specific for a nuclear factor present only in non-seed tissues. This factor would determine gene expression in embryonal tissues only. The importance of CAAT and TATA sequences in the control of site-specificity has also been proved.

Four Soya nuclear factors that interact with specific sequences present in the  $\alpha^\prime$  and  $\beta$  URS of the gene have been identified as well. Two of those embryonal factors, SEF3 and SEF4, bind to sites inside the enhancer region (from -257 to -79 ). SEF3 binds to the middle of the sequence 5'-AACCCA.....AACCCA-3', exclusively in the gene encoding  $\alpha'$  subunit. the Accumulation and degradation of this protein (SEF3) is paralleled by accumulation and degradation of lpha' mRNA, supporting the hypothesis that SEF3 be involved in the control of  $\alpha^\prime$  expression. As compared to SEF3, SEF4 is present in a lesser amount, has many recognition sites (5'-A/GTTTTA/G-3') in  $\alpha'$ , but mostly in  $\beta$ . The presence of this factor is correlated to the regulation of  $\boldsymbol{\beta}$ expression (Lessard et al., 1991). Deletion and sitespecific mutagenesis experiments have proved that the sole action of these factors does not affect the sitespecificity nor the expression level, coordination with

the activity of the other regulatory elements being necessary. However, on the basis of the homology with light-induced proteins, these proteins are supposed to have a regulating role only under certain conditions (Fujiwara and Beachy, 1994).

Embryonal factors with a behavior similar to the SEF3, as verified in gel shift experiments, have been found in tobacco as well (Lessard et al., 1991). This and other data obtained with GUS activity assays under control of Soya  $\alpha'$  promoter prove that the site- and stage-specific control mechanism is conserved (Lessard et 1991; Riggs et al., 1989). Ιt has also been binding hypothesized that tobacco trans factors 35SCaMV promoter may also interact with the legumin boxes (Katagiri et al., 1989). None of the afore mentioned factors appears to be directly responsible of the time regulation, and no NRS-like factor has been found possessing a negative control as in the case of bean (Bustos et al. 1991).

More recent studies, concerning some legumins and vicilins in Vicia Faba, contradict some generalizations on the regulation of the storage proteins expression in seeds (Wobus et al., 1995), showing that expression of genes of B4 and LeB4 Vicia Faba legumins is not limited to embryonal tissues, nor are they temporally restricted to the cell expansion phase in embryogenesis. Proteins are stored for short periods of time and then degraded in all embryonal tissues, suspensor and endosperm included, within well-defined developmental stages. This is probably in order to allow an uninterrupted supply to the embryo of compounds having a high C and N content during all the growth and differentiation stages. Therefore, this data allow to hypothesize that the seed proteins expression be also controlled metabolically, and not merely at a developmental stages level. The possible relationship existing between storage accumulation and carbohydrate metabolism (soluble glycid

level) is presently being investigated. Since all classes of seed storage proteins share a similar behavior in the different species, this data require a careful evaluation of the behavior, in terms of expression, also for the Soya proteins  $\beta$ -conglycinine and 7S basic globulin. Data resulting from the study, which the present invention is based on, clearly show the tissutal specificity of the expression of the structural portion of lactoferrin under control of both the promoters. Instead, the activation phase was at the present not investigated in detail as the sole capability of total seed accumulation was of interest. Specifically concerning the 7S basic globulin, it is a storage protein of the Soya seed, with a high methionine and cysteine content. Alike β-conglycinine, also 7S basic globulin (Bg) is stored in seed in large amounts (3% of seed total proteins). It consists of two subunits, one of 27 KDa, the other of 16 KDa encoded by same mRNA, linked by disulfide bridges. synthesized as sole precursor polypeptide consisting of a two subunits. putative peptide signal and of polypeptide is processed to yield the mature dimeric protein. In the genome about four copies of the Bg gene are present (Watanabe and Hirano, 1994).

This protein is mainly located in the seed embryonal tissues and its expression pattern is unusual for a storage protein. In fact, a portion of Bg is accumulated in the intercellular spaces of the cotyledon parenchyma (Nishizawa et al., 1994), whereas at an intracellular level it is stored in protein bodies on the middle lamella of cell wall and in the plasma membrane (Watanabe and Hirano, 1994). This location suggests that the Bg is not a mere storage protein, having other functions as well. More accurate data on Bg location and expression period in Soya are not available. It has never been verified whether the site- and time-specific expression mechanism be preserved in other transformed vegetal species (like tobacco). To this end, reference is made to

general data on storage proteins and to studies on Bg homologues in lupine (conglutyn  $\gamma$ ), with which it has a high sequence homology. This protein is stored only in lupine embryonal tissues (cotyledons and embryonal axis) 40 days after blooming. It has not been detected in other tissues such as leaves and sprouts. In seeds of transgenic tobacco, the conglutyn  $\gamma$  gene is increasingly expressed from the 15th to the 20th day after blooming until the 40th, then begins to decrease (Ilgoutz et al., 1997).

One of the peculiar features of the Bg is that it is secreted in large amount from Soya seeds soaked in water at 40-60°C. It is uncertain whether the secreted proteins are neosynthetized after heat-treatment, or instead are the proteins already present to be secreted. Since a post-heating increase in specific mRNA has been highlighted, it is assumed that the Bg is actually synthesized as a consequence of the thermal shock (Hirano et al., 1989).

Not much is known on the regulation mechanism of the of the gene encoding the Βq protein, nevertheless sequences in the promoter region involved in the gene regulation have been identified. Besides the CAAT and TATA box sequences, respectively located at -116 and -25 with respect to the transcription start site, three regulatory elements similar to thermo-specific sequence enhancers present at the non transcribed 5' region of genes in other organisms, have been detected. These heat shock elements (HSE) consist of two 5 pb conserved units: 5'-NGAAN-3' and 5'-NTTCN-3'. thermoregulated promoter of the Soya heat-shock protein, the enhancer elements, observed also in Bg as well, cooperate synergistically with three CCAAT box sequences located upstream thus increasing gene expression; these putative sequences are present also in the bg promoter.

Sequences responsible of the site- and timespecificity expression were not identified. The interest for this protein derives from the fact that it is accumulated in high amounts in the Soya seed (3% of total proteins) and therefore has a strong seed-specific promoter which can ensure a high level of expression of the gene it controls. Moreover, it is known that the regulation mechanism of this protein is different from that of the other storage proteins of Soya seed but the details are not known. However, studies on the promoter and on its site- and time-specific activation mechanism have never been carried out using reporter genes in transgenic plants.

CONG, as storage proteins, Both Вq and synthesized exclusively in the seed tissue and are stored in large amounts in cells constituting this organ, inside specific compartments. Concerning the transcriptional and above all the post translational regulation level, it runs through the mechanisms of transport intracellular and of compartimentalization, which are to date to be clarified in many aspects.

In fact, those mechanisms involve all processes influencing the concentration, retention and distribution of the proteins in the endomembrane system (Okita, 1996).

However, general principles of protein targeting do exist, valid for all plant species.

- 1. Targeting information are contained in the proteins themselves, as discrete signals. Those signals are intercepted by specific recognition signals such as receptors or simple interactions with membrane lipids.
- 2. Different types of signals do exist (topogenic sequences) each with a specific function. Among them there are signal sequences that start the protein translocation across specific endomembranes and interact with receptors/translocators facilitating the unidirectional transfer. Then there are stop and retention sequences that block the transfer to the membrane or to the inside of the compartment. Selection

sequences target proteins to the various cellular compartments. All those elements can be of a sequential type, i.e. localized in the N-terminus, central or C-terminus portion of the protein, or conformational, i.e. consisting of amino acids which although nonsequential, are yet adjacent in the native protein conformation. Moreover, there may be various signals simultaneously, and they can be modified or activated (e.g. by phosphorylation). After transfer the signal is often deleted using specific cleavage sites for endogenous proteases.

mRNA accumulation in a particular region influences the protein location. Soya seed storage proteins, globulins as well as lectins, are stored in storage vacuoles. In fact, several types of vacuoles do exist. Some of them, besides having the function of maintaining the turgor pressure and of regulating ion, sugars and amino acid release, also constitute the depository of storage and defense proteins. The specific signal sequence for targeting to the vacuole has not been identified yet, apart from some plant species (Kermode, 1996). Probably, one or more surface regions of the correctly conformed protein are recognized selection mechanism. Plant cells possess the unique feature of accumulating storage proteins in the protein bodies, whereas in animals similar inclusion bodies are formed only when an excess of protein synthesis occurs. Therefore, the latter protein bodies consist of unmuddled accumulation of conformationally incorrect or partially The formation processed proteins. and organization process of the protein bodies in plants remains unclear, although it is known that it consists of a series of ordered events (Okita, 1996). Globulins are proteins with an acidic isoelectric point (pI), accordingly they are translocated in the endoplasmic reticulum (ER) and in the Golgi complex as soluble proteins. As soon as they reach the vacuoles, due to their low pH or possibly to the

processing and assembly, these proteins precipitate forming particled aggregates that will thereafter originate the protein bodies (Kermode, 1996).

In leguminosae, different storage proteins are accumulated in the same protein bodies with no spatial segregation. In other plant species the protein bodies form in the ER and are then absorbed in vacuoles by autophagocytosis (Kermode, 1996).

is well-grounded general pattern βconglycinine though the specific as well, targeting sequence have not been identified for this protein. Instead, the  $\beta$ -conglycinine binding with a BIPhomologous protein has been observed. This protein functions as chaperonine and, just like other proteins in different plant species, can have the role of retaining  $\beta$ -conglycinine in the ER until its correct conformation is reached (Galili et al., 1993; Shewry et al., 1995; Kermode, 1996; Fontes et al., 1996). As for 7S basic globulin, available information is scarce. It is known that it is located in protein bodies on the middle .lamella of cell wall and in plasma membrane and not in vacuoles (Watanabe, 1994). For this reason, its mechanism of division into compartments is hypothesized to be different from that of  $\beta$ -conglycinine. However, it is known that even wall-located proteins follow the same transport pathway of the vacuole proteins, translated in the ER, then transported to Golgi and lastly secreted to the outside or inserted the membrane by vesicular traffic.

However, storage proteins expression in heterologous hosts shows that the compartimentalization mechanism is universal. In transgenic plants the seed vacuole storage proteins are correctly targeted. Nevertheless, sometimes transport can be inefficient, especially in vegetative organs with respect to seeds. In tobacco, leguminose storage proteins are correctly targeted to vacuoles both in seeds and in leaves, yet in leaves there is a lower

accumulation level. This is so because a difference exists in the transportation efficiency or because of a different processing rate (different proteases or higher instability). Hence, it can be understood how seed-specific in plant production of a heterologous human protein is a complex mechanism, so that the preliminary verifying of the functioning and efficiency of the expression system, as constructed in a model host organism, constitutes an important step.

It has been seen how tobacco is one of the most widely used plants to this end. Its preferential use in assays derives from the fact that it is one of the better known plants, both in a genetic and in a biological and physiological respect. This, together with the ease of effecting the genetic transformation and the shortness of vegetative cycle, made it become an important model for biotechnological experiments, a model transformation specific systems and micropropagation conditions are now better known. Additionally, tobacco further advantage of a near-complete possesses the extensibility of the obtained results to several other plant species, consequence of the high conservation rate of genic control mechanisms, that precisely proved to be usually highly conserved in other plant species, and in particular in leguminosae. Therefore it is particularly for the study of the promoters taken out suitable and in particular of their capacity of therefrom, allowing a gene of interest to be expressed in a controlled way in a transformed plant. Genes of interest are usually those encoding proteins suitable alimentary pharmaceutical or field. Accordingly, heterologous gene of interest for this kind application is that of the human lactoferrin, a protein belonging to the transferrin family, and as such capable to stably and reversibly bind two iron ions.

In fact, by virtue of its biological functions lactoferrin turns out to be interesting from a

nutritional as well as from a pharmacological point of view. It is present in human milk and has a fundamental role in neonatal feeding, as a matter of fact several have been attributed · biological functions protein, among which a bactericidal and bacteriostatic against wide range of а microorganisms and the capacity of increasing iron absorption at the intestinal level (Lonnerdal and Iyer, Lonnerdal, 1993). Hambraeus and Moreover promotes cellular growth, controls myelopoiesis and is modulating the inflammatory capable of response (Lonnerdal and Iyer, 1995; Oguchi et al. 1995; Penco et al. 1995).

Therefore, at first, attempts to research in milk of other mammals a protein capable of binding iron and possessing the same properties were carried out.

It has been observed that milk of all mammals contains two types of iron-binding proteins, present in identical different ratios: transferrin, to milk transferrin, and lactoferrin. Human has particularly high lactoferrin content, in fact concentration in colostrum is of 5-10 mg/ml, although it decreases during lactation to about 1 mg/ml in ripe milk (Hambraeus and Lonnerdal, 1993). However, the amount of lactoferrin is much lower in milk of other species, like goat, horse, pig and mouse, In cow's milk for instance its concentration is of about 0.1 mg/ml. In some species such as rabbit, rat and dog, lactoferrin is absent, the prevailing iron-binding protein being instead transferrin.

Further, lactoferrin (LF) produced by other nonhuman mammal species, assumes in each of them different structural characteristics, and therefore different properties.

Human lactoferrin (LFU) is a 78 KDa monomer glycoprotein, with a bilobate structure. There is a high degree of homology between the N-terminus domain and the

C-terminus one, both at the amino acidic sequence (37%) and at the tridimensional structure level. tridimensional structure has been described in detail by X-ray crystallography (Lonnerdal and Iyer, 1995). The gene encoding LFU has been cloned and sequenced. Genic control mechanisms at a transcriptional and translational level and estrogens and iron role in those mechanisms are also known (Liu et al, 1991). The mature protein consists of a 692 aa polypeptidic chain with a 8.8 - 9 pl. It contains 16 disulfide bridges and shows some resistance 1995), has three glycanic to proteolysis (Lommerdal, polyacetylactosaminic chains bound with N-glycosidic bonds to the amino acidic residues Asn233, Asn476 and Asn545 and the molecular weight of the glycosilated protein is 82 KDa.

One of the most important differences existing among lactoferrins (LFs) present in the various animal species is precisely the glycosidic chain composition. In fact, unlike human LF, bovine LF contains  $\alpha-1,3$  galattosidic residues and glycans of oligomannosidic type; the role of the glycosidic chains has not been defined yet, however it is possible that glycans protect LFU against attacks from proteolytic enzymes. Each of the two LFU domains is capable of binding tightly, yet reversibly to a ferric ion and at the same time to a carbonate or bicarbonate ion molecule (Hambraeus and Lonnerdal, 1993). binding sites in human milklactoferrin completely saturated, but only at 6-8% of their capacity (Stowell et al., 1991).

In recent years, a series of studies aimed at understanding the mechanisms of action and the relation between molecular structure and function of this protein have been carried out. The strategy adopted was that of studying LF molecules structurally altered by site-specific mutagenesis. Accordingly, expression systems of LFU recombinants producing in a simple and inexpensive way a protein as identical as possible to the one

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purified from human milk were carried out.

However, all heterologous hosts used so far for the LFU recombinant are eukaryotes, as, it being a complex glycoprotein, requires a sophisticated processing apparatus.

1991 Stowell et al. cloned the LFU gene in cultured neonate hamster renal cells. An inducible  $2n^{2+}$ promoter and the secretion signal of a hamster endogenous protein were used to maximize expression. Concentration of LFU recombinant secreted in the culture medium was of about 20 mg/l, sufficient for crystallization therefore for structural studies. Characterization revealed that it has the same molecular mass of native LFU maintaining intact the iron-binding site. It only differs from human milk LFU in the glycosidic chains and N-terminus sequence, but these do not influence folding. Such an expression system is highly expensive and not suitable for the production of the amounts of proteins required at an industrial level.

Then LFU was expressed in transgenic mice's milk. In this case the entire animal rather than the cultured cells was transformed, using the regulation sequence of the bovine gene  $\alpha$ -Sl caseine. It was shown that LFU mRNA is exclusively expressed in female mammary gland during lactation. In milk the protein reaches a  $0.1-36~\mu g/1$ concentration. Recently this LFU recombinant has been characterized (Nuijens, 1997), and it has been observed that it has the same molecular mass, N-terminus sequence and immunoreactivity of native LFU. It also maintains the capacity of releasing iron at acidic pH and the bond to bacterial lipopolysaccharides. Also in this case, glycosilation, as well as in vivo bactericidal antiinflammatory action, are different from the ones in native LFU.

Indeed, a highly significant production system of recombinant LFU is that carried out in Aspergillus awamori (Ward et al, 1995). This method, which is

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patented, yields commercial amounts of recombinant LFU (2 g/l). In order to maximize expression LFU is produced as a fusion protein with part of the structural gene, regulation sequence and secretion signal in the culture medium of the glucomylase. The fusion polypeptide is processed to yield mature LFU by an endogenous peptidase. Glucoamylase is an Aspergillus protein expressed in high amounts.

An alternative eukaryotic host is the one used by Mitra and coworkers in 1994. They have transformed tobacco cells in suspension. In the transgenic calluses a protein much shorter than the native LFU and therefore found to be unstable is produced in small amounts. Recombinant LFU shows activity against phytopathogenic bacteria, e.g., Xantomonas campestri, Pseudomonas syringae and others. In the above-mentioned study the obtaining of entire and fertile plants is not reported.

Recently LFU was also produced in culture in insect cells, using Baculovirus as expression system (Salmon et al., 1997). This is a highly powerful expression system, yielding a recombinant protein identical to the native one, apart from the glycosilation level. Nevertheless it maintains the binding with the specific receptors.

All the above reported expression systems allow to obtain an amount of protein sufficing for functional studies, and in some cases (Aspergillus) for commercial uses as well. In the latter case however, safe use of the purified protein, e.g. in milk substitutes for neonates, requires an excessive purification in order to ensure the absence of immunogenic or allergenic substances. However, transgenic plants have never been used to this end, nor products directly suitable in human nutrition have been ever yielded, for instance by the recombinant LFU expression in alimentary plants.

#### Summary of the invention

The present invention relates to a general system of tissue-specific, and in particular seed-specific,

recreatively that all most in the

accumulation of heterologous proteins, designed and carried out with the object of maximizing the production while preventing the degradation thereof, by using leader sequences and promoters of Bg and  $\beta$  conglycinine genes. To this end, the structural part of the selected gene may encode proteins having an enzymatic activity, used in human therapy or in industrial processes, or proteins with a general (lactoferrin) or specific (antibodies or antigens) pharmacological activity, or antibody proteins for phytopurification or for the elimination of mycotoxins present in foods.

The present invention also relates in particular to a system that, enabling the in plant tissue-specific expression of the human lactoferrin gene, provides an important solution to the problem of the production of this protein. This system in fact determines the production of plants capable of expressing relatively high amounts of this protein that, in the preferred embodiment providing the expression of a synthetic gene designed by the inventors so as to maximize its in plant expression, reaches industrially relevant levels. Moreover, such transgenic plants allow to avoid the costly product purification processes, as they can be used as nutraceuticals, and therefore being directly intaken as alimentary products. Accordingly, also the use for the production of protein flours or of protein extracts yielded from tissues, and specifically from seeds of the afore mentioned transgenic plants, for the production of functional foods or special preparations for children is possible.

The protein accumulation in the seed further allows to significantly increase the iron content of the same, or of the flours derived therefrom. Accordingly this system allows the obtaining of foods which, having a twofold iron content with respect to analogous product, can be also used for supplying this micronutrient's deficiency through a normal alimentation.

These plants can anyhow be used also for human lactoferrin purification, by conventional methods based on chromatography techniques.

This expression system further provides the use of new recombinant vectors, constructed by testing the effect of various leader sequences and processing sites of the mature protein, enabling the production of any protein of interest and in particular of lactoferrin or of fragments derived thereof, with a tissue-specific protein accumulation, in plants belonging to different families among which leguminosae, cereals, solanaceae, fruit-bearing plants and horticultural produce general. In particular they are structured so as to have the following functionally linked components: promoter; (b) a signal sequence; (c) a nucleotide sequence optimized for in plant expression corresponding to the amino acidic sequence of the entire lactoferrin or to fragments thereof; human (d) polyadenylation signal.

In particular the case concerning plasmid is considered, and wherein regulation elements and signal sequences used are those belonging to two genes encoding storage proteins that are very common in Soya seeds, i.e. a  $\beta$ -conglycinine and a 7S basic globulin, isolated and cloned from the Richland soya variety. They can be used to transform plant cells both by the Agrobacterium method and by direct physical methods (Gelvin and Schilpercort, 1995; DuPont Biolistic Manual, DuPont). The vegetal transformed cells are hence selected with the selection agent provided for the purpose and induced to form entire fertile plants capable of forming seeds, in turn capable of expressing the gene for lactoferrin, and accumulating it as storage protein.

This result was obtained by designing and synthesizing an artificial gene encoding the same amino acidic sequence of the natural human gene, but having sequence mutations such that codons most frequently used

by the human cell are replaced with those most frequently used by the vegetal cell. This result yielded the change of the 31% of the codons in the original gene (see table 1). The remarkable production of human lactoferrin detected in the various plants transformed with the synthetic gene, with yields going from 1% to 1.8% of the seed total storage proteins, but not in plants having the natural gene, proved the in plant functionality thereof. Furthermore on the basis of such sequence, which allows the efficient expression of the human lactoferrin in any vegetal cell, a person skilled in the art can derive the specific sequence expressed with a higher efficiency in each single vegetal species, by the simple application of the common general knowledge.

Concerning all the above disclosed, object of the present invention is a polynucleotide encoding human lactoferrin, characterized in that it has a sequence totally or partially corresponding to the sequence reported as SEQ ID NO:1, or to a sequence biologically equivalent thereof and in that said sequence is optimized for in plant expression, and in particular the case wherein said polynucleotide has fused to the 5'-terminus end a sequence selected from the group comprising the sequences shown as SEQ ID NO: 13 and NO: 14. A particular case of sequence partially corresponding to the SEQ ID NO:1 is given by a sequence corresponding to one or more fragments of said sequence.

Object of the present invention is also the human lactoferrin protein, obtained from the expression of the afore mentioned sequences.

A further object of the present invention is a recombinant DNA vector, in particular a plasmid, comprising at least one sequence of a gene of interest, in particular the gene encoding the complete human lactoferrin, specifically a sequence totally or partially corresponding to the SEQ ID NO:1, operatively linked to regulation elements enabling the tissue-specific

expression of said gene. A special case is the one where such regulation elements consist of an expression cassette for plants allowing tissue-specific expression of said gene.

Further cases of interest are those wherein the expression cassette for plants are constituted by the regulation elements of the protein 7S basic globulin, and in particular when among said regulation elements there is the sequence reported as SEQ ID NO:21, or the regulation elements of the  $\beta$ -conglycinine protein, and in particular when among said regulation elements there is the sequence reported as SEQ ID NO:22, the case wherein the sequence of the gene encoding complete or partial human lactoferrin is operatively linked, or even fused, to a leader sequence, and the case wherein such leader sequence is selected from the group comprising sequences SEQ ID NO: 13 and NO: 14.

Of particular importance is the case wherein such plasmid is selected in the group comprising vectors pBI, pGEM or pUC.

further object of the present invention Α constituted by the transformation process of vegetal cells wherein the transformation is effected with one of the above-mentioned vectors, the transgenic vegetal cells can be obtained through transformation of wild type vegetal cells with at least one of the same vectors, and by the cells that anyhow contain a gene of interest, or portions thereof, and in particular the gene encoding human lactoferrin operatively linked in an expression cassette, enabling the tissue specific expression of the gene itself, in particular that of the gene encoding 7S globulin and that of the gene encoding basic conglycinine.

A further object of the present invention is constituted by cellular aggregates and in particular calluses characterized in that they are obtainable by the aforesaid cells.

A further object of the present invention are also the transgenic plants obtainable from the aforesaid cells with conventional techniques, or anyhow containing a gene of interest and in particular the gene encoding human lactoferrin, specifically the one with a sequence corresponding totally or partially to SEQ ID NO:1, operatively linked in an expression cassette enabling the tissue specific expression of the gene itself.

Of particular relevance is the case wherein such transgenic plants are selected from the group comprising solanaceae, cereals, leguminosae, horticultural produce and fruit-bearing plants in general, in particular Soya, tobacco and rice, wherein the gene encoding lactoferrin is specifically expressed in the storage tissues or in the fruit. A further object of the present invention is the use of such transgenic plants as nutriceuticals.

A further object of the present invention is constituted by the production processes of functional foods containing proteins produced by the aforesaid transgenic plants, of vegetal milks, starting from natural and/or concentrated proteins deriving from the above-mentioned plants, and anyhow any human lactoferrin production process, characterized in that it utilizes the aforesaid plants.

Lastly, object of the present invention is also the human lactoferrin obtained from the aforesaid transgenic plants.

The invention will be better described with the aid of the annexed figures.

## Description of the figures

Figure 1 shows the strategy adopted for the assembly of the synthetic gene encoding human lactoferrin.

Figure 2 shows the map of plasmids pGEM-PGLOB(A) and pGEM-PCONG (B) obtained from the cloning of the Soya promoters in plasmid pGEM (Promega), in which the restriction sites used to derive the plasmids are highlighted.

Figure 3 shows agarose gel electrophoresis analysis of digestion of plasmids pGEM-PGLOB with Sal I (lanes 1, 2, 3 and 4) and pGEM-PCONG CON Sph I (lanes 6, 7, 8 and 9), carried out to test clockwise orientation of the insert. All PGLOB samples tested positive, yielding fragments of the expected sizes. PGLOB 1 is sample 1 selected for the subsequent molecular work. In contrast, PCONG samples did not yield the expected pattern, suggesting the possibility of the presence of errors due to the adopted cloning technique (hypothesis later discarded, see figure 6) or due to isolation of a variant of the control region that in the Richland variety differs from the one of the disclosed sequence (Dare variety). Sample 5 is lambda marker HindIII.

Figure 4 shows agarose gel electrophoretic analysis of the restriction pattern of two plasmid pGEM-PCONG clones with enzymes Nde I (lanes 1 and 2), Rsa I (lanes 3 and 4), and SnaB I (lanes 9 and 10), carried out in order to test orientation and identity of the constructs. Cuts with Nde I, Apa I and SnaB I yielded the expected patterns in contrast to the cut performed with Rsa I; this results are justified from the differences found in sequence and reported in figure 6. In lanes 5 and 6 markers of molecular weight  $\lambda$ -DNA Hind III and Marker IV (Boehringer) respectively are present.

Figure 5 shows electrophoretic analysis of the restriction of various clones of plasmid pGEM-PCONG with Rsa I (lanes 1-6) and Hinf I (lanes 9-14) enzymes in order to test identity of constructs. In both cases the obtained pattern do not mirror the expected ones, but are conserved among the different clones, thereby suggesting their being due to differences in the original sequence with respect to the published restriction map and not to errors in the amplification phase with Taq polymerase or in the cloning. Adopted markers are  $\lambda$ -DNA Hind III and Marker IV.

Figure 6 shows the comparison between the published

sequence of gene CONG promoter region and the one cloned in plasmid pGEM-PCONG.

Figure 7 shows a schematic view of the two plasmids resulting from the cloning of the native gene LFU into the two vectors pGEM-T and pBI121, carried out in order to obtain plasmids used later on as transformation control.

Figure 8 shows the map of the two plasmids resulting from synthetic LFU gene cloning into vectors pGEM-PGLOB and pGEM-PCONG, i.e. plasmids pGEM-PGLOB-LFU (A) and pGEM-PCONG-LFU (B), respectively.

Figure 9 shows the map of plasmids pBI-PGLOB-LFU (A) and pBI-PCONG-LFU (B) wherein the restriction sites used are highlighted. In particular, box (A) shows the construction of a plasmid containing the synthetic gene represented in the sequences list as SEQ ID NO:1 and cloned in plasmid pBI101 fused to promoter PGLOB and in Open Reading Frame with the "leader" of 7S basic globulin.

In contrast, box (B) shows the construction of a plasmid containing the synthetic gene reported in the sequence list as SEQ ID NO: 1 and cloned in plasmid pBI101 fused to promoter PCONG and in open reading frame with the  $\beta$ -conglycinine leader.

Figure 10 shows electrophoretic analysis of the restriction of various clones of plasmid pBI-PCONG-LFU with enzymes Xba I, BamH I, Sac I; samples 4 and 5 test positive. In position 6 molecular weight marker Ladder 1Kb is present.

Figure 11 shows electrophoretic analysis of the restriction of two clones of plasmid pBI-PCONG-LFU with enzymes Sal I (lanes 1 and 2) and with Xba I and Sac I (lanes 4 and 5); Both samples tested positive. Samples 3 and 6 represent the positive controls, i.e. pGEM-PCONG-LFU digested with Sal I, Xba I and Sac I respectively.

Figure 12 shows agarose gel electrophoresis analysis of PCR products from genomic DNA extracted from various

plants transformed with pBI-PGLOB-LFU, performed using primers PLT48 and PLT49 for the promoter sequence PGLOB. Positive samples 2, 3, 4 and 5 represent the band of the DNA amplified to 1500 base pairs, while samples 6, 7 and 8 represent the negative control of PCR and the positive control (pGEM-PGLOB) respectively. Molecular weight markers Ladder 1Kb are found at 1 and 9.

Figure 13 shows in box (A) the agarose gel with the genomic DNA of tobacco transformed with pBI-PCONG-LFU (lanes 1-5) or with pBI-PGLOB-LFU (lanes 6-9) cut by enzyme BamH I. M is the molecular weight marker Ladder 1 Kb. Sample 10 shows the positive control pGEM-LFU, not shown in the photo for quantitative reasons. In box (B) the hybridization pattern of human LF on the genomic DNA of the same tobacco plants is shown; samples 1, 2 and 3, belonging to plants PCONG 1, PCONG 3, and PCONG 4 respectively are positive, as is the case for samples 5, 7 and 8, belonging to plants PGLOB 10, PGLOB 3 and PGLOB 4 respectively. It is evident that pGEM-LFU, the positive control (lane 10), was only partially digested as also the super-coiled plasmid forms are present.

Figure 14 shows SDS-PAGE electrophoretic analysis of proteins partially purified from seeds of the transgenic plants tested with Southern analysis of the preceding figure (A) and \_Western analysis of the same proteins, after transfer to a membrane, using polyclonal antibodies specific for the human lactoferrin (B). In particular, in electrophoretic analysis of xod (A) SDS-PAGE (30 from mature seeds cellular proteins DAP) transgenic tobacco is shown. In position 2, 3, 4 and 5, 7 the same samples tested positive to Southern analysis, extracted by buffer at pH 2.7 and pH 7.6 respectively, are found. Samples 8 and 10 represent the (milk-extracted human lactoferrin, control positive Sigma) and the negative control (non-transformed plant of the same variety), while in position 9 the molecular weight marker Rainbow (Amersham) is found. In box (B)

autoradiography of anti-lactoferrin antibody hybridization with the same proteins transferred to DEAE-nitrocellulose membrane is shown; the sample corresponding to plant PGLOB 10, in position 2 and 5, does not yield a positive signal, although according to Southern analysis it is transformed. All other samples are positive.

Figure 15 shows electrophoretic analysis of extracted from seeds and leaves transgenic tobacco transformed with the plasmids of which at figure 9. In particular, in box (A) protein coloration carried out with Coomassie blue is reported. In box (B) Western Blotting carried out with human LFU-specific antibodies on the proteins of the gel shown in box (A) after transfer to membrane is reported. In particular, in lane 1 plant PGLOB 1, with leaf-extracted proteins is reported, in lane 2 always plant PGLOB 1, with seedextracted proteins is reported, in lane 3 plant PGLOB 3, leaf proteins, is reported, in lane 4 plant PGLOB 3, seed proteins, is reported, in lane 5 plant PCONG 105, leaf proteins, is reported, in lane 6 plant PCONG 105, seed proteins is reported, in lane 7 plant PCONG 105, seed proteins treated with N-deglycosilase F, is reported (see text), in lane 8 the molecular weight marker is reported, in lane 9 the human LFU present on the market treated with N-deglycosilase F is reported, in lane present on the market is reported.

Figure 16 shows Western analysis of LFU protein extracted from human milk and of recombinant protein isolated from tobacco seed, before and after N-deglycosilase F enzyme treatment. Analysis was performed with human lactoferrin specific antibodies. In lane 1 LFU extracted and purified by HPLC from seeds of plant PCONG 105 is reported; in lane 2 a protein as in 1 after a 18-hours treatment with N-deglycosilase F is reported; in lane 3 commercial LFU after a 18-hours treatment with N-deglycosilase F is reported; in lane 4 commercial LFU is

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reported, a diminution is apparent in the molecular weight of the two enzyme-treated samples (2 and 3).

## Detailed description of the invention

strategy adopted for the generation transgenic plants capable of producing human lactoferrin was developed along two directions: on the one hand, expression comparative analyses on plant systems, particularly tobacco and Soya, have been carried out, in order to have a basis for the designing of a sequence encoding human lactoferrin thereof, sequence optimized to maximize its expression in vegetals. Accordingly in the sequence designing the necessity that the required posttranslational modifications for the production in the active form could be effected on the translated protein, and that, both for its conformation and due to subcellular localization, the protein be sufficiently stable to be accumulated in relevant amounts in the transformed plants, was taken into account. This proved having ascertained after various crucial, carried out in the past years the impossibility of an in plant production of human lactoferrin using constitutive expression systems (e.g. promoter 35S) as well promoters inducible by leaves cut. Moreover, besides difficulties related to the type of promoter used, the production level and the stability of the protein were depending be scarce and on preferential use of the codons between the human gene and the plants.

Therefore, a plasmid vector system was developed containing newly synthesized utilizing vectors a lactoferrin gene regulated by tissue- and stage-specific promoters capable of yielding a high gene expression and of accumulating the protein in a stable and efficient way inside seed storage organs. Moreover the selection of leader sequences and the design of the fusion point between those and the structural portion of the mature yielded lactoferrin protein that, in protein a

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quantitative and possibly also in qualitative terms, has the same glycosilation level and the same amino terminal sequence of the native protein, which is important for some of its functional characteristics.

Concerning the synthetic gene design, all the necessary and possible triplets were modified taking into account their preferential use in the two reference plants, tobacco and Soya. In particular, data represented in table 1 were used.

TABLE 1

COD	0NS	HUNAN	SOYA	TOBACCO	म्बर्ग सर	LTU SYN
A.P.G	CZ A	5.5	4.5	4-8	3	3
	<b>ದಾರ</b>	11.2	7.5	3.8	5	5
	CCC	10.7	2.2	2.2	7	
			7.4	7.7		3
	ಞಾ∣	4.5				8
	AGA	9.8	14.8	12.5	13	16
	AGG	10.5	11.3	12.1	13	<u>10</u>
LEU	CIA	6.1	7.0	7.4	0	7
المست	CIC	20.1	16.2	12.7	20	1
		42.1	10.3	7.1	28	9 8
	CIG					8
	CIT	10.8	23.7	21.8	9	19
	TTA	5.4	8.4	10.3	2	2
	TIG	11.1	20.5	20.7	9	18
527	TCA	9.7	14.7	15.8	5	. 11
2-7		17.B	9.5	10.0	13	
	TCC					. 7
	TOTAL	4-1	4.3	4.3	, 2	2
	TCT	13.3	17.3	20.6	10	, 14
	AGC	18.7	16.8	9.0	13	9
	AGT	9.9	14.0	11.9	В	· B
TER	ACA	14.3	14.9	16.4	8	В
J.F.Pt				11.9	. 9	,
	ACC	22.6	14.0			7
	ACG	. 6.6	3.2	4.0	2	- 2
	ACT	12.5	17.4	.20_4	12	14
33C	CA.	15.4	. 30.6	25.2	6	1.2
	<u> </u>	20.6	10.5	9.0	10	5
	CCG	6.8	4.6	3.2	5	4
		16.1	22.2	21.7	. 13	14
					12	18
ALA	GCA.	14.4	20:5	22-3		10
	GCC	29.7	16.6	15.2	25	16
	GCG	7_2	4.7	4-4	' В	В
	ecr	18.9	23.2	35.21	18	21
CT.Y	GGA	• 17.4	22.7	31.4	11	21
	GGC	25.3	11.9	14.0	24	В
				10.0	12	. B
	ccc	17.5	11.0			36
	CCI	11.5	22.1	30.0	5	16
VAL	GIA	6.1	7.5	11.5	2	3
	GTC	16.2	9.0	13.8	7	5
	ere i	30.7	25.8	14.4	22	18
	GII	10.2	24-4	29.4	5	15
TVC					<del></del>	22
LYS	AAA	21.5	23.2	23.2	22	
	AAG	35.2	35.2	20-71	23	23
スニム	AAC	22.3	29.2	25-0	17	17
	AAT	16.5	20.2	27.8	14	14
GIN	CAA	10-E j	27.4	22.5	8	. 14
·				· · · · · · · · · · · · · · · · · · ·		
	ದ್ಯಾ	33.2	20.7	14.1	21	15
HIS	C3C	14.7	€.5	8.9	4	4
	CAF	9.3	12.1	11-17	5	5
GLU	GAA	25.4	34.5	25.7	15	12
	GAG	41.5	25.8	25.2	21	15
224	GAC	28.5	18.8	17.2	22	16
	GRT	21.5	29.7	33.0	16	22_
TV	TAC			16.6	12	12
TYR		18.0	17.1			
	TAT	12.3	. 15.2	20.5	9	9
C.72.	TGC	13.8	9.9	8.1	וֹדֹנ	17
•	767	9.5	3.5	10-1	15	15
PHE						
F.T.2	TIC	22.1	23.5	18-1	17	15
	TII	15.8	19.3	25.2	10	12
ILI	AIA	6.11	12.2	10.71	4	- 4
	ATC	24.3	1	13.7	7	5
	>77	25.0	15.0			7
			23.71	29.4	5	<del></del>
<u> </u>	ATG	22.31	20.34	22.51	4.1	
<del></del> 2	ا تئ:	11.7!	22.51	13.71	30 (	10
					_ <u> </u>	

In carrying out such operation, the value G+C and A+T of the two systems (human and vegetal), the non-tandem repeat of some triplets that may cause shifts in reading, etc. were also taken into due account.

Synthetic LFU gene was then obtained using primers reported in the annexed sequence listing from SEQ ID NO:8 to SEQ ID NO:12 and from sequence SEQ ID NO: 15 to SEQ ID NO: 20 and following the assembling strategy reported in figure 1, consisting in repeated PCR cycles, using for each cycle different pairs of synthetic primers allowing the gradual elongation and the forming of the final sequence as designed. Similarly on the basis of such final sequence the sequences contained the codons preferably expressed in various species of interest, among which rice, have been obtained.

parallel, also native LFU gene (wild type) encoding human lactoferrin was cloned, always by PCR technique, starting from a cDNA library of human mammalian tissue (Clontech). The gene was recovered in its structural part lacking the signal peptide and the poly-A site and cloned in pGEM-T to form plasmid pGEM-LFU whose map is represented in figure 7. Primers designed for amplification are reported in the annexed sequence listing at sequences SEQ ID NO:2 and SEQ ID NO:3; those added the restriction site BamHI 5' at restriction site SacI at 3'. After checking the sequence, which tested identical to the published one, the yielded natural gene was cloned in vector pBI121, on sites BamHI and SacI under control of promoter 35S (see figure 7), this plasmid (pBI-LFU) was then denominated pBI-35S-LFU and used as control in the genetic transformation experiments and the subsequent molecular biochemical analysis.

Concerning the preparation of the recombinant vectors containing the elements that allow the tissue-specific expression of the LFU gene in expression cassettes for plants, we proceeded as follows. In order

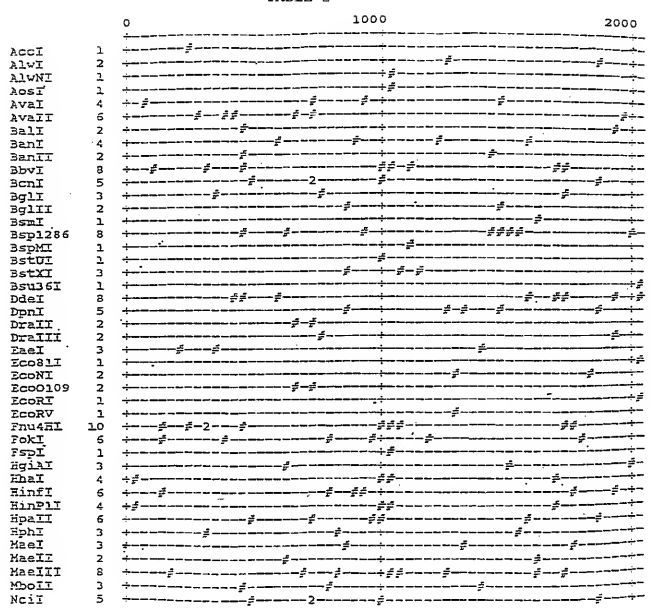
to obtain the seed-specific expression of the protein the promoters and signal sequences of two genes encoding storage proteins that are very abundant in Soya seeds, i.e. a  $\beta$ -conglycinine (CONG) and a 75 basic globulin (GLOB) were used.

These regulation sequences were isolated and cloned from Soya, Richland variety.

In particular, to clone the two GLOB and CONG sequences PCR technique (PCR = Polimerase Chain Reaction; Innis et al. 1990) was used. In this case genomic DNA extracted from Soya leaves of Richland cultivar was used. Oligonucleotides used for specific amplification are reported in the annexed sequence listing from SEQ ID NO: 4 to SEQ ID NO: 7.

For the GLOB promoter the cloned region includes the entire regulation sequence and the sequence encoding the signal peptide (leader) plus the first codon of the structural sequence, such sequence is indicated with SEQ ID NO: 13 in the annexed sequence listing. For the CONG promoter the cloned region includes the entire regulation sequence and the sequence encoding the signal peptide plus the first codon of the structural sequence, such sequence is indicated with SEQ ID NO: 14 in the annexed sequence listing. For both regulation sequences the most suitable restriction site for insertion proved to be XbaI (TCTAGA), while downstream proved to be BamHI (GGATCC), both absent from the native and synthetic lactoferrin sequence, as highlighted in the following tables 2 and 3.

TABLE 2



## TABLE 2 (continues)

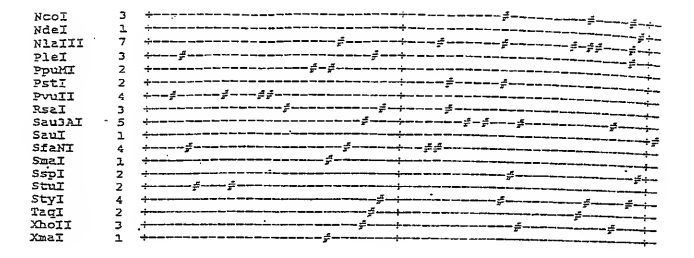


TABLE 3

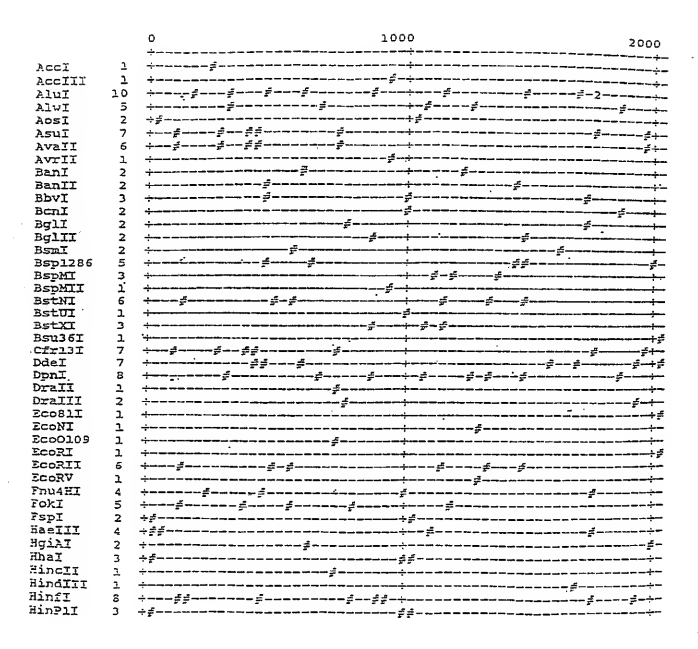
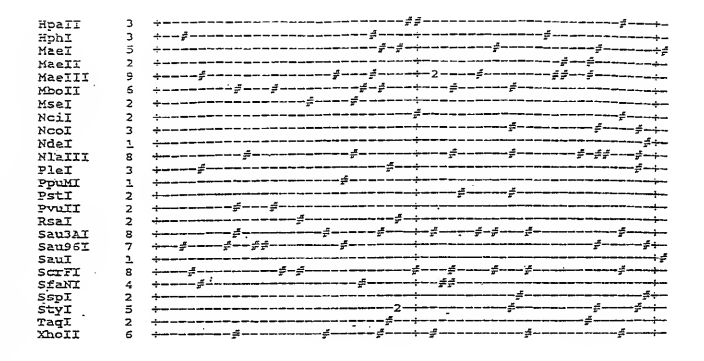


TABLE 3 (continues)



DNA template was extracted from Glycine max leaves, Richland variety, and amplification products match sizes expected for GLOB (1515 pb) and CONG (1163) promoter on the basis of EMBL sequence data.

Therefore, starting from fragments amplified by ligation in vector pGEM-T, the two vectors pGEM-PGLOB and pGEM-PCONG, whose map is reported in figure 2, were constructed. Yielded plasmids were tested by restriction analysis performed with several enzymes chosen among those cleaving in a limited number and with an overall sequence distribution (see figures 3, 4 and 5) and a clone for each type was selected and sequenced. Sequenced clones showed to be significantly different from the expected sequence. As an example, a comparison between the data bank promoter CONG sequence and the one obtained sequencing clone pGEM-pCONG is reported in figure 6. A 5% difference was detected, therefore the two promoters can be considered as different.

The synthetic gene for human lactoferrin was cloned at first in plasmids pGEM-PGLOB and pGEM-PCONG, cut with enzymes BamHI-SacI, to form plasmids pGEM-PGLOB-LFU and pGEM-PCONG-LFU respectively, whose map is disclosed in figure 8, and then the construct XbaI-SacI transferred in vector pBI101 cut with the same enzymes. In the event of a plant transformation carried out with physical means, as for rice in our case, plasmids pGEM-PGLOB-LFU and pGEM-PCONG-LFU, can be directly used after addition of a terminator, in cotransformation with a vector containing the selection marker (e.g., a PUC-type vector containing the gene for hygromycin resistance). Resulting plasmids pBI-PGLOB-LFU and pBI-PCONG-LFU, whose map is reported in figure 9, were used in the genetic transformation of the after an accurate control carried restriction with various enzymes to assay the correct integration of the DNA construct (figures 10 and 11).Plasmids pBI-35S-LFU, pBI-PGLOB-LFU and pBI-PCONG-LFU were transferred in A. tumefaciens EHAlO5 strain cells,

made competent by electrophoresis. Strains containing the three plasmids were used to transform about 450 leaf disks (LD) of tobacco, Petit Avana variety. Formation first of shoots and then of roots was induced from calluses formed on leaf disks (LD) in presence of kanamycin. Once rooted, plants were potted and at least 50 kanamycin-resistant plants were analyzed for each construct. The same plasmids containing, this time, the hygromicyn-resistance genes marker, are used also for the transformation of rice. In this case in particular 40 plants hygromicyn resistant were analyzed.

Plants  $T_0$ , of rice as well as of tobacco, were tested by PCR technique (figure 12), assaying the presence of the lactoferrin gene inside the genome of the tested plants; plants  $T_1$  were assayed by Southern analysis (figure 13), that compared to PCR technique allows a more accurate testing of the transgene presence in the genome, and with Western analysis (figure 14) allowing detection of genic product and therefore the functionality of inserted gene.

All plants with native LFU gene under control of promoter 35S led to accumulation of a protein, recognized by LFU-specific antibodies, of a molecular weight lower than 50 KDa. This protein was found in small amounts in leaves, becoming undetectable in the developed leaves. Plants of rice as well as of tobacco transformed with the two constructs pBI-PGLOB-LFU and pBI-PCONG-LFU produce and accumulate exclusively in seed protein having a molecular weight corresponding to the glycosilated human protein as shown by electrophoretic analysis of extracted proteins and by the related Western Blotting carried out with LFUspecific antibodies (see figure 15). Presence recombinant protein exclusively in the seed and not in the leaves was assayed in all the examined transgenic plants (about 50 for the two constructs) with Western techniques.

Recombinant LFU protein isolated from seed and purified with HPLC technique showed to be identical to the native protein concerning its iron binding capacity and its inhibiting effect towards the examined bacterial strains. Treatment with a deglycosilating enzyme confirms the presence of posttranslational modifications in all alike, at present at least in quantitative terms, to those present in native lactoferrin as highlighted by Western analysis, the results of which are disclosed in figure 16.

Moreover, the contribution of iron consequence of the introduction of the lactoferrin gene in rice plants was assessed. In particular, in the following table 4 results in terms of iron content of some transgenic lines of the Ariete and Rosa Marchetti varieties transformed with plasmids pBI-PCONG-LFU and pBI-PGLOB-LFU are reported. Iron content was measured by atomic absorption after flour mineralization with HNO3 e H2O2.

Table 4: Analysis of iron content in Ariete (A) and Rosa Marchetti (RM) varieties and of the respectuive transgenic lines capable of in-seed accumulation of protein lactoferrin.

Sample	Fe Concentration (µg/g)	RSD
Ariete	34.3	3.0
A cl. 2-3	48.5	0.7
A cl. 5-1	86.5	1.8
A cl. 6-2	117.0	1.2
A cl. 6-3	72.4	3.2
Rosa Marchetti	23.5	3.6
RM cl. 5-3	64.5	2.8
RM cl. 2-5	76.5	2.4
RM cl. 3-6	52.7	3.0
RM cl. 4-1	48.2	1.7

It is therefore evident from all of the abovereported results that using the native gene, described in the literature, for human lactoferrin under control of the traditional promoters used for genetic transformation of plants, human protein lactoferrin cannot be produced in relevant amounts, in a stable form and with the posttranslational modifications typical of this protein.

So far a general description has been given of the present invention. With the aid of the following examples, a more detailed description will now be given of specific embodiments thereof, with the purpose of giving a clearer understanding of objects, features, advantages and methods of application of the invention.

EXAMPLE 1:

# Agrobacterium Tumefaciens-mediated tobacco transformation Day 1

small amount of Agrobacterium tumefaciens strain EHA 105, taken from a petri plate culture with a sterile loop so as not to exceed in the amount thereby avoiding subsequent problems in controlling bacterial proliferation on plated leaf disks, was inoculated in 2 ml of sterile LB. Then, from a healthy tobacco plant of Avana variety a leaf showing no alteration whatsoever, conversely showing optimal turgor condtions, was taken. The leaf was briefly washed in bidistilled water to remove surface impurities, immersed for 8 min in a 20% sodium hypochlorite and 0.1 % SDS solution and left to dry under a vertical flow hood. From then on all steps were carried out under hood. In particular, the leaf was immersed in 95 ક ethanol and shaken in order to completely wet the pages thereof (letting the petiole emerge) for 30 - 40 sec. The leaf was then allowed to dry out completely.

Disks were obtained from the entire leaf surface with an ethanol-sterilized punch, let fall on plates with MS10 free of antibiotics; in particular, the ratio of 30 disks per plate was not exceeded.

Next, 2 ml LB + (just inoculated) Agrobacterium were poured on plate, and the bacterial suspension was evenly spread over the entire plate with a gentle rotatory movement, in order to obtain an homogeneous bacterial

distribution among the disks. LB in excess was carefully aspirated with a pipette. In the course of those steps at all times a parallel negative control was provided by means of a plate to which nothing, or only LB was added.

Then plates were incubated at 28°C for 24-48 hours in constant lighting conditions, and bacterial growth was indicated by the appearance of a thin opaque layer spreading over the entire plate.

Day 2

Leaf disks (=LD) were carefully transferred on a plate with MS10 + 500 mg/l cefotaxime, and incubated at 28°C for 6 days in constant lighting conditions. This step determines the Agrobacterium inactivation.

Day 8

LD were then carefully transferred on a plate with MS10 + 500 mg/l cefotaxime and 200 mg/l Kanamycin, and incubated at 28°C for 14 days in constant lighting conditions. This step determined a selection of the transformed plants: in fact, gene of kanamycin resistance was carried by the plasmid inserted in Agrobacterium.

Day 22

LD, that in the meantime had grown developing a callus, were carefully transferred on a plate with MS10 + 500 mg/l cefotaxime, 200 mg/l Kanamycin and 500 mg/l carbenicillin, and incubated for 6 days. This step determines elimination of the agrobacteria possibly survived to the previous antibiotic treatments (a very frequent occurrence).

Day 28

LD were transferred again on MS10 + 500 mg/l cefotaxime and 200 mg/l Kanamycin, and incubated until shooting. When shoots showed at least two leaves, they were separated from the callous mass and transferred on the radication medium: MSO + 500 mg/l cefotaxime and 200 mg/l Kanamycin.

At the appearance of roots, seedlings were extracted from the plate, freed from agar residues, gently washed

in running water and planted out in loam and sand (2:1) inside small plastic pots. Soil was previously saturated with water, then pots were covered with transparent plastic lids to preserve high humidity conditions, and placed in a growth chamber at room temperature, with a daily 16-hour lighting period.

#### EXAMPLE 2:

## Rice transformation by physical methods

Rice seeds of Ariete and Rosa Marchetti varieties were harvested at milky ripening, when the endosperm is still liquid. The embryo was isolated with a lancet after removal of the two glumes. Immature embryos are of different size and shape, depending on the number of days elapsed from blooming: The ones deemed most suitable for the in vitro culture and the successive transformation, i.e. those of an 1.5 mm average size, were cultured on a medium containing 2,4-D auxine to promote scutellum cell division and suppress differentiation of young embryos, obtaining cellular proliferation of the scutellum area. embryos this stage underwent bioholistic transformation, with the following parameters: particle size 1.5-3 μm, particle concentration 500 μg, membrane rupturing pressure 1.100 psi, membrane-microcarrier gap 6 mm, microcarrier stop-point gap 6 mm, stop-point target gap 10.5 cm.

Transformation was effected by a cotransformation technique, using as a selectable marker the gene for plasmid pROB5. Cotransformation on hygromycin effected with a total DNA concentration of 1  $\mu g/\mu l$  using for bombing and with а DNA plasmid/suitable plasmid (pROB5 with pBI-PGLOB-LTU or with pBI-CONG-LTU) ratio of 1:1 assessed on the number of molecules (abt. 1:4 in amount).

Osmotic conditions of the plant material were optimized carrying out a preculture on 3% saccharose and plasmolysis prior to bombing on MS with 10% saccharose for 1 hour. 24 hours after bombing with PDS-1000/He

bioholistic system the material was transferred on 3% saccharose medium.

For selecting the transformational events, bombed tissue underwent selection in presence of hygromycin B (Duchefa). 1 day after bombing embryos were transferred on solid MS medium, additioned with 2 mg/l 2,4-D auxine, 50 mg/l cefotaxime, 50 mg/l hygromycin B, 3% saccharose 0.35% agarose. One week later embryos transferred on R2 liquid medium (Ohira et al., containing 1 mg/l thiamine, 50 mg/l cefotaxime, 50 mg/l hygromycin B and 3% saccharose, pH 5.8, in 190 ml plastic vessels (Greiner). Liquid cultures were shaken at 90 rpm and 28°C in the dark, replacing the liquid medium every 7 days. After 3-4 weeks resistant calluses formed on the embryo surfaces are transferred on R2 solid medium additioned with MS vitamins, 2 mg/l 2,4-D auxine, 50 mg/l cefotaxime, 50 mg/l hygromycin B, 60 g/l saccharose and agarose and maintained for 2 - 4weeks, formation of embryogenic structures.

At the formation of embryogenic structures calluses were transferred on solid MS regeneration medium containing 2 mg/l BAP, 0.2 mg/l NAA, 3% maltose, 50 mg/l hygromycin B, 50 mg/l cefotaxime and 0.8% agarose.

Embryogenic calluses were maintained in phytotron at 28°C with 16 hours of lighting to induce formation of shoots, that, once formed, were transferred on hormone-free 1/2 MS radication medium, 3% saccharose and 0.3% gelrite. After 3-4 weeks at 28°C, seedlings were transferred in a Yoshida solution (Yoshida et al. 1976) and maintained at 25/19°C day/night with a 11 h day length. After 4 weeks plants were potted and grown in hothouse until the cycle end.

#### EXAMPLE 3:

Purification of lactoferrin protein from different tissues of the plant and assessment of molecular weight.

Extraction of all the proteins of tobacco seed was performed grinding the seeds in liquid nitrogen in

presence of an extraction buffer (0.5 M saccharose, 0.1% ascorbic acid, 0.1% Cys-HCl, 0.01 M Tris-HCl, 0.05M EDTA pH 8).

Then the solution was centrifuged for 30 minutes at  $14.000~\rm rpm$  at  $4\,^{\circ}\rm C$  and the supernatant was kept with the soluble proteins.

Then the solution was filtered with filters of 0.2  $\mu$ m porosity, and the lactoferrin partially purified by removing proteins of a molecular weight lower than 30 KDa by centrifugation in Centricon 30 column (Amicon).

The lactoferrin was further purified by HPLC chromatography on Resource Q column (Pharmacia) at a weak cationic exchange, with elution in phosphate buffer pH 7 and NaCl gradient 20-100%. The peak corresponding to lactoferrin eluted at 0.7 M NaCl.

The fractions of the elution range were reunited and filtered in Centricon 30 to remove salt.

For the lactoferrin extraction from tobacco leaves, up to the centrifugation step we proceeded as in the case of extraction from seed, then the supernatant was additioned with 60%  $(NH_4)_2SO_4$  and left shaking in ice for 60 min.

Then the solution was centrifuged at 14.000 rpm for 15 minutes at  $4^{\circ}\text{C}$ , the pellet recovered and then suspended again in phosphate buffer pH 6.8.

For the assessment of molecular weight in SDS-PAGE, the colorant (SDS loading buffer) was additioned to the lactoferrin sample (20µl) and the samples were loaded onto 8% polyacrilamide minigels. Running conditions were: initially 10mA, and 20 mA for the entire run, in Trisglycine lx buffer. Then the gel was stained by Silver staining technique and the molecular weight assessed referring to molecular weight standards.

EXAMPLE 4:

Western analysis of the lactoferrin protein produced in plant and deglycosilation thereof.

Lactoferrin purified from seed according to example

2, after electrophoretic separation on acrylamide gel was transferred by electroblotting (buffer 25mM Tris, 192 mM glycine, 20% methanol, 45 V at  $4^{\circ}$ C) to a nitrocellulose membrane (BA85 Schleicher and Schull).

The membrane with the immobilized protein was shaken for 60 min in TBS-T 5% Skin milk solution and then, after some washings, with the same solution containing the primary antibody in a 1:2500 ratio.

After reaction with primary antibody the membrane was washed and placed in contact with the secondary antibody (Anti-Rabbit peroxidase conjugate), always in TBS-T Skin milk solution, in a 1:12.000 ratio.

After reaction with secondary antibody the membrane was washed several times and placed in contact with Amersham's chemiluminescence kit ECL.

The membrane was then exposed in contact with a photoplate (Hyperfilm MP, Amersham) in darkroom for variable lengths of time.

Deglycosilation with N-glycosidase F enzyme (Boehringer Man.) was carried out using 10  $\mu$ l in volume of glycopeptide (10  $\mu$ g) denatured in 0.1% SDS brought to boiling point for 2 min.

To this solution 90  $\mu$ l of buffer (20 mM phosphate buffer pH 7.2, 50 mM EDTA pH 8, 10 mM sodium azide, 0.5& NP40, 1%  $\beta$ -mercaptoethanol) were additioned and it was brought to boiling point again for 2 min, then cooled at 37°C.

To the resulting 100  $\mu$ l 1 U of N glycosidase F was additioned and let incubate at 37°C for 18 hours. Then the reaction product was analyzed on SDS-PAGE gel and the lactoferrin protein detected by Western technique.

#### GLOSSARY

The term "recombinant polynucleotide", as it is used here to characterize a polynucleotide useful in the production of lactoferrin, relates to a polynucleotide of genomic origin, cDNA, semi-synthetic or synthetic, that, by virtue of its origin or manipulation: 1) is not

associated to a portion or to the totality of the polynucleotide to which it is associated in nature, and/or 2) is linked to a polynucleotide differing from that to which it is associated in nature, or that 3) does not exist in nature.

The term "polynucleotide", as it is used here, relates to a polymeric form of nucleotides of any length, ribonucleotides as well as deoxyribonucleotides. This term exclusively refers to the molecule primary structure. Hence, the term includes single and double stranded DNA as well as single and double stranded RNA. It also includes modified forms of the polynucleotide, e.g. by methylation, phosphorilation or "capping", and non modified forms.

An «expression cassette for plants» relates to a recombinant polynucleotidic sequence obtained by linking together operatively various elements constituted by the polynucleotidic sequences that determine the in plant expression of a character and that are easily transferable as discrete constructs, from a vector to another by enzymatic restriction.

A "vector" is a replicon to which another polynucleotidic fragment is added, in order to effect the replication and/or expression of the fragment itself.

A "replicon" is any genetic element, for instance a plasmid, a chromosome, a virus, that behaves as an autonomous polynucleotidic replication unit inside a cell; therefore it can replicate autonomously.

"regulation sequence" refers to polynucleotidic sequences that are needed to effect the expression and/or the secretion of coding sequences to which they are bound. The nature of these regulation sequences differs depending on the host; in prokaryotes those regulation sequences usually include promoter, binding site of ribosomes and in eukaryotes these regulation sequences usually include promoters, terminators and, in some cases, enhancers. In addition, in prokaryotes as well as eukaryotes, leader sequences control the host cell

secretion of the expressed polypeptide. The term "regulation sequences" includes, at least, all components whose presence is required for expression, and may also include additional components whose presence is advantageous, for instance leader sequences.

A «leader» sequence is a polynucleotidic fragment, usually short, encoding a transport signal of the protein fused thereto and leading the protein transfer into specific cellular compartments. If the transfer takes place through the endoplasmic reticulum the protein undergoes specific posttranscriptional modifications.

"Operatively linked" relates to a juxtaposition wherein the above described components are in a relation enabling them to function in the expected way. A regulation sequence «operatively linked» to a coding sequence is linked in such a way that the coding sequence expression takes place in conditions that are compatible with the regulation sequences.

AN open reading frame, ORF is a polynucleotidic sequence region encoding a polypeptide; this region can represent a portion of coding sequence or a complete coding sequence.

A "coding sequence" is a polynucleotidic sequence that is transcripted in the mRNA and/or translated in the polypeptide when placed under control of appropriate regulation sequences. The ends of the coding sequence are determined by a translation start codon at 5' and by a translation stop codon at 3'. A coding sequence can include, without being limited to, mRNA, cDNA, and recombinant polynucleotidic sequences.

"Recombinant host cells", "host cells", "cells", "cell lines", "cell cultures" and other terms indicating microorganisms or cell lines of superior eukaryotes, cultivated as unicellular entities, are used here in an interchangeable way. They relate to cells that can be, or have been, used as hosts for recombinant vectors or other transfer polynucleotides, including the progeny of the

cell that was originally transformed. It is implicit that, due to random or deliberate mutations, the progeny of a single parental cell need not necessarily be identical to the parental cell from a morphological and a genetic point of view. Progenies of the parental cell that are sufficiently similar to the ancestor cell and can be characterized for their salient capacity, as e.g., the presence of a nucleotidic sequence encoding the peptide of interest, are included in the progeny understood according to this definition and fall within the same terms.

For «cell aggregation» a group of cells that are not structured in an organized tissue, but result from an undifferentiated proliferation of cells maintained in particular conditions of hormonal concentration.

"Transformation", as it is used here, refers to the insertion of a exogenous polynucleotide in a host cell, regardless of the method used for the insertion itself, e.g. direct acquisition, Agrobacterium infection, sexual reproduction. The exogenous polynucleotide can be maintained as a non integrated vector, for example a plasmid or, alternatively, it can integrate in the host genome.

As it is used here, the term "polypeptide" relates to the amino acidic product of a sequence encoded inside a genome and does not relate to the specific length of the product: accordingly, peptides, oligopeptides and proteins are included in the definition «polypeptide» This term does not relate to the post-expressional modifications of the peptide, as e.g. glycosilation, acetylation, phosphorilation, sialilation and the like.

A "wild type polypeptide", has an amino acidic sequence identical to the one encoded in the genome of the organism source of the coding sequence.

"Native lactoferrin" and analogous terms relate to the lactoferrin isolated from the source in which it is usually produced in nature by a genome existing in nature.

A "non-native polypeptide" refers to a polypeptide

that is produced in a host differing from the one wherein it is produced in nature.

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# CLAIMS

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- 1. A polynucleotide coding for human lactoferrin, characterized in that it has a sequence totally or partially corresponding to the sequence reported in the annexed sequence listing as SEQ ID NO:1, said sequence being optimized for the in plant expression.
- 2. The polynucleotide according to claim 1, wherein said polynucleotide has fused at the 5'-terminus the leader sequence of 7s basic globulin gene or the leader sequence of  $\beta$ -conglycinine gene
- 3. The polynucleotide according to claim 2, wherein said leader sequence is the sequence reported in the annexed sequence listing as SEQ ID NO: 13 or the sequence reported in the annexed sequence listing as SEQ ID NO: 14.
- comprising DNA vector recombinant 4. A polynucleotide according to any of claims 1 to 3 operatively linked to regulation elements allowing the expression of said polynucleotide.
- 5. The recombinant DNA vector according to claim 4, wherein said regulation elements is a plant expression cassette allowing the tissue specific expression of said polynucleotide.
  - 6. The vector according to claim 5, wherein said plant expression cassette includes the promoter of the gene coding for the protein basic globulin 7 S.
  - 7. The vector according to claim 6, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:21
- 8. The vector according to claim 6 or 7, wherein 30 said plant expression cassette includes the leader sequence of the gene coding for the protein basic globulin 7 S.
- 9. The vector according to claim 8, wherein said leader sequence is the sequence reported in the annexed 35 sequence listing as SEQ ID NO: 13.

AMENDED SHEET

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- 10. The vector according to claim 5, wherein said plant expression cassette includes the promoter of the gene coding for  $\beta$ -conglycinine protein.
- 11. The vector according to claim 10, wherein said . promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:22.
  - 12. The vector according to claim 10 or 11, wherein said plant expression cassette includes the leader sequence of the gene coding for the  $\beta$ -conglycinine protein.
  - 13. The vector according to claim 12, wherein said leader sequence is the sequence reported in the annexed sequence listing as SEQ ID NO: 14.
  - 14. A vegetal cell including the polynucleotide according to any of claims 1 to 3.
  - 15. A vegetal cell including the vector according to any one of the claims 4 to 13.
  - 16. A cellular aggregation obtainable from cells according to claim 14 or 15.
- 17. The cellular aggregation according to claim 16, 20 wherein said aggregations are calluses capable of regenerating transgenic plants.
  - 18. A transgenic plant including in a tissue cell the polynucleotide according to any of claims 1 to 3.
  - 19. The transgenic plant according to claim 18, wherein said tissue cell is a storage tissue cell.
    - 20. The transgenic plant according to claim 18, wherein said tissue cell is a fruit tissue cell.
- 21. The transgenic plant according to any of claims 18 to 20, said plants being selected from the group 30 consisting of solanaceae, cereals, leguminosae, fruit bearing plants and horticultural plants.
  - 22. The transgenic plant according to claim 21, said plant being selected from the group consisting of Soya, tobacco and rice.
  - 23. Use of the vector according to any of claims 4 to 13 for the transformation of vegetal cells.

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- 24. Use of the transgenic plant according to any one of the claims 18 to 22, for the production of nutriceuticals.
- 25. Use of the transgenic plant according to any of claims 18 to 22, for the production of human lactoferrin.
  - 26. Use of the transgenic plant according to any one of claim 18 to 22, for the production of lactoferrin flours or of lactoferrin extracts obtained from tissues of said transgenic plant.
- 10 27. Use of the transgenic plant according to any one of the claims 18 to 22 for the production of functional foods containing lactoferrin.
  - 28. The use according to claim 27, wherein said functional food is selected from the group consisting of vegetal milks, fruit juices, fruit and/or vegetable homogenized foods.
  - 29. A plant expression cassette allowing the tissue specific expression of a gene of interest comprising the promoter of the gene coding for the protein basic globulin 75.
  - 30. The plant expression cassette according to claim 29, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:21
  - 31. The plant expression cassette according to claim 29 or 30, wherein said plant expression cassette includes the leader sequence of the gene coding for the protein basic globulin 7 S.
    - 32. The plant expression cassette according to claim 31, wherein said leader sequence is the sequence reported as SEQ ID NO: 13.
    - 33. A recombinant DNA vector comprising a gene of interest under the control of the plant expression cassette according to any of claims 29 to 32.
  - 34. The vector according to claim 33 when depending on claim 31 or 32, wherein said gene of interest is fused to the leader sequence.

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- 35. A plant expression cassette allowing the tissue specific expression of a gene of interest comprising the promoter of the gene coding for the  $\beta$ -conglycinine protein.
- 36. The plant expression cassette according to claim 35, wherein said promoter has the sequence reported in the annexed sequence listing as SEQ ID NO:22
  - 37. The plant expression cassette according to claim 35 or 36, wherein said plant expression cassette includes the leader sequence of the gene coding for the leader sequence of the gene coding for the  $\beta$ -conglycinine sequence of the gene coding for the  $\beta$ -conglycinine protein.
  - protein.

    38. The plant expression cassette according to claim

    37. wherein said leader sequence is the sequence reported

    as SEO ID NO: 14.
  - as SEQ ID NO: 14.

    39. A recombinant DNA vector comprising a gene of interest under the control of the plant expression cassette according to any of claims 35 to 38.
- 40. The vector according to claim 39 when depending on claim 37 or 38, wherein said gene of interest is fused to the leader sequence.
  - 41. A vegetal cell including the vector according to any one of the claims 33, 34, 39 or 40.
  - 42. A cellular aggregation obtainable from the cell according to claim 41.
  - 43. The cellular aggregation according to claim 42, said aggregations being calluses capable of regenerating transgenic plants.
  - transgenic plants.

    44. A transgenic plant including in a tissue cell
    the vector according to any of claims 33, 34, 39 or 40.
  - 45. The transgenic plant according to claim 44, wherein said tissue cell is a storage tissue cell.
  - 46. The transgenic plant according to claim 44, wherein said tissue cells is a fruit tissue cell.
  - 47. The transgenic plant according to any of claim 43 to 45, said plants being selected from the group

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consisting of solanaceae, cereals, leguminosae, fruit bearing plants and horticultural plants.

- 48. The transgenic plant according to claim 47, said plant being selected from the group consisting of Soya, tobacco and rice.
- 49. Use of the vectors according to claims 33, 34, 39 or 40 for the transformation of vegetal cells.

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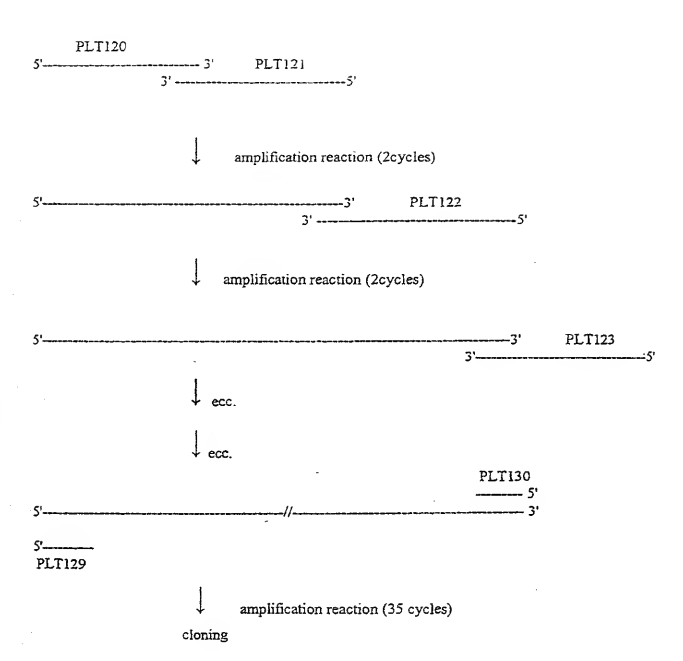
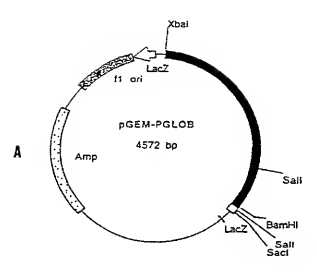


FIG.1



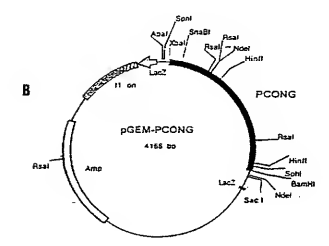


FIG.2

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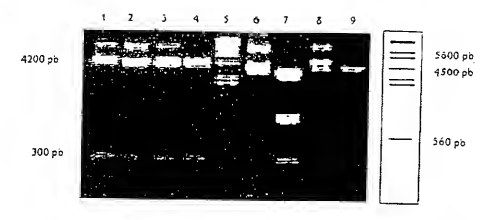


FIG.3



FIG.4

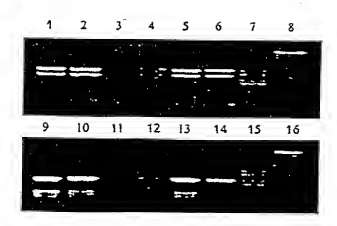
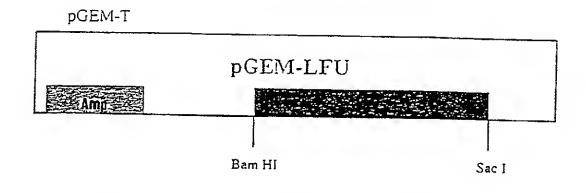


FIG.5

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gmapsp	TETAGASTTTTEAAATTTSAATTTTAATSTSTSTTGTAAGTATAAATTTAAAATAAAAAT
PCO::GL7Sp6	TOTAGASTTTTEAAATTTTAATTTTTATSTSTSTATAAAATTTAAAATAAAAAT
9=Ээsр	ANAACAATTATTATATCAANATIGCAAAA-CATTTAATAGITATTATTATTAATAAAA
РСЭ::377\$p6	AAAACAATTATTATATCAAAATSGCAAAAATATTTAATACCTATTATTTAAGAAAAAA
gm±psp ?CONST7Sp5	TATGTAATAATATATTATATTITAATATCIATTCTIATGTATTTTTAAAAATCTATTA TATGTAATAATAATATATTATAT
gmbpsp COMGUT7Sp6	TATATTTATCAACTAAAATATTTTTATATCTACACTTATTOCATTTTATTAATTTTC TATATTTATCAACTAAAATATTTTTATATCTACACTTATTGCATTTTTATCAACATTTT
çmipsp PCC::GT7Sp€	TTS:STTTTTTGCCATAT.TTAATA-TGACTATTCTTTAATAATGACTATTATTATTATAATAATCATTATTATTATTATTATTA
9720952	TOOTACATATTOTTOGAACCATATGAAGTOTTCATTSCATTTGACTATGTGGATAGTGTT
2021:377596	TCGTAGATATTGATGGAACCATATGAAGTGTCCATTGGATTCGACTATGTGGATAGTGTT
g=bpsp 2CO:IGT7Sp4	THEATTCATGCCCYTCATTTOSCECT - ATTAATTAATTTASTAACAGATTCCTTCTAATC
9m2psp	AGTTAGTTAATGETTGETTGATGATAATTTAATET-GGTAGT-TGGAATGEGATAATATTGA
70012077Sp6	AGTTAGTTATTTTCCTGCATGATAATTAATGTTGGTAGTGGAATGGTAGAACACTGA
çabpaş PCCHCT7Sp5	TTAGTTTTTGACACAAAAAAAAAAAAAAAAAAAAAAAAA
gmopsp	GTAT-COTT-TOCATAGCAATGTTTAAGTTCATAAAATTCAAACAAAAACGCAATTAAACAC
PCONGT7Sp6	GTATCCTTTGCATAGCAATGTCTAAGTTCATAAAATTCAAACAAA
empsp	AGTOG-ACATCA TITATCCA-CTAGCTGATCAGGATCGCCCCCCCCAAGAAAAAAAAACCTG
PCONGT7Sp6	AGTGGGACATCACTTATCCACTAGCTGATCAGGATCGCCCCCCTCAAGAAAAAAAA
gmbpsp	GACTERAAAGETATTERACAACAAGAETTACTEREAAAGETSTEAATEGAGE-GETTAAA
PCOHUTTSpi	GEACTERAAAGETATTEREAACACACTATTEREAAAGSTETEAATETAACAAGETAAA
ფი <b>ბ</b> ენე 7€00 <b>::07</b> 75ე-6	ACATTCACCAACTCAACTCATCATCATCACCACACATTTCTTC
- <del>peomost</del> ebe Gwebeb	ARTESTATTITECTTCCSCCAC-TCATSTITTATTTCAALACCCSTCAAACTSCATCCAACTCCATCAACTCCACTCC
5201:311254	CACCCCUTTOCCCAAATSTT CATGCATUTTAACAASACTTATGACTATAAATATCTUTAAT
642522	CACCCCGTGGCCAAATGTCCATGCATGTTAACAAGACTTAIGACTATAAATATATCTGCAAT
gmapsp	CTCSGCCCAAGTTTTCATCATCAAGAACCAGTTCAATATCCTAGTACGCCGTATTAAAGA
FCCVCT7Sp6	CTCGGCTGAGGTTTTGATCAAGAACCAGTTCAATATCCTAGTACAGGGTATTAAAGA
STOPSF PCCMCT7Sp4	ATTTAAGATATACTATGATGAGAGUGGGUTTCCCATTACTGTTGCGGGGGTGGTTTTACCATTTAAGATACAACTACTATGAGAACACCAGTTCCCATTACTGTTGCTGG-AGTTGCTTGCCAGTTACTGTTGCTGG-AGTTGCTTGCCAGTTACTGTTGCTGG-AGTTGCTTGCCAGTTACTGTTGCTGG-AGTTGCTTGCCAGTTACTGTTGCTGG-AGTTGCTTGCTGG-AGTTGCTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTTGCTGG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-AGTG-
gebpsp	TAGEAT DAS. TTETSTETSTETTTSGCATTGCSTATTSSSAAAASCA-AACCCCASTCACA
PCDWGT7Sp6	TGGCATCASTTTCTGTTTCATTTGGCATTGSGTATTZSSAAAAGCAGAACCCCAALTCACA
çebeş	ACAMOTOCOTOCOAAGTTOCAATAGCGAGAAAGACTTTTAGAGGAACGAACGAAGGATGCAAGG
FCC::CTTSp&	ACAACTSCTTCCGAAGTTSGAAATGCGAGAA-GACTTCTACAGGAACCAAGGATGCCAAG
gröpsp	CTCSTTSCAACCTCCTTAAGGTSGGATCT
PIC:IGT7Sp4	CTCSTTSCAACCTCCTTAAGGTSGGATCC

FIG.6



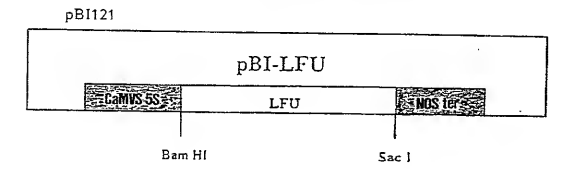
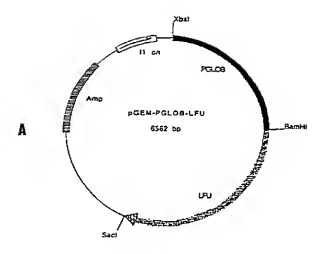


FIG.7

PCT/IT99/00226



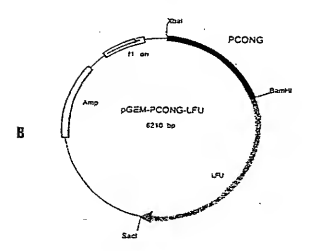


FIG.8

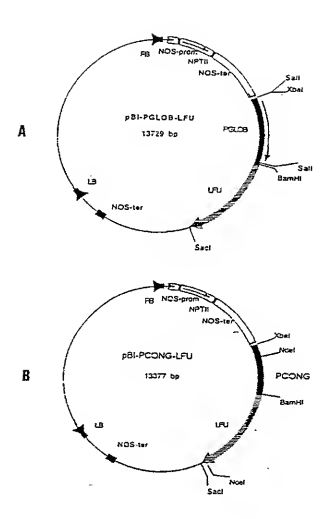


FIG.9

8/13

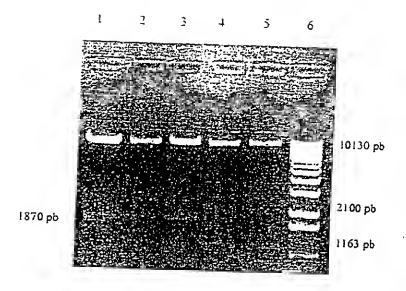


FIG. 10

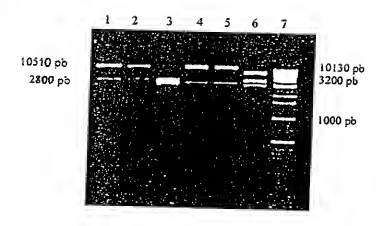
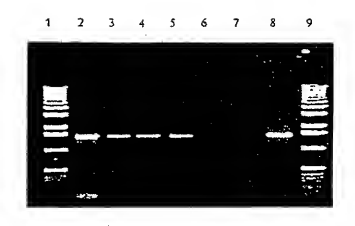


FIG. 11



**EIG.12** 

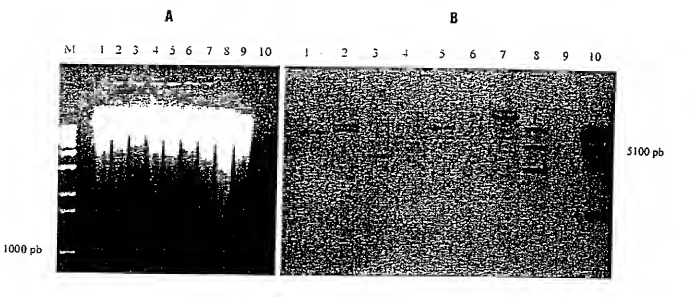


FIG.13

82 kDa

B
1 2 3 4 5 6 7 8 9 10 1 2 5 4 5 6 7 8 9
82 kDa

FIG.14

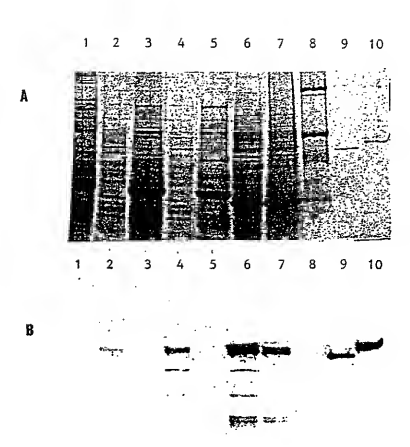


FIG.15



FIG. 16

· 8	PATENT
Attorney's D	ocket No. <u>B-4075PCT_618484-4</u>
(ORIGINAL, DE	COMBINED DECLARATION AND POWER OF ATTORNEY SIGN, NATIONAL STAGE OF PCT, SUPPLEMENTAL, DIVISIONAL, CONTINUATION, OR CIP.
As a below n	amed inventor, I hereby declare that:
[ ] [ ]	TYPE OF DECLARATION  ion is of the following type: (check ona applicable item below)  original  design
NOTE: If the continuation one of last the	supplemental he declaration is for an International Application being filed as a divisional, or continuation-in-part application, do <u>not</u> check next item; check appropriate brae items. national stage of PCT
NOTE: If one DIVISIONAL, [ ]	e of the following 3 items apply, then complete and also attach ADDED PAGES FOR CONTINUATION, OR CIP.  divisional continuation continuation-in-part (CIP)
WARNING:	INVENTORSHIP IDENTIFICATION  If the inventors are aach not the inventors of all the claims an explanation of the facts, including the ownership of all the claims at the time the last claimed invention was made, should be submitted.
believe I am original, first	e, post office address and citizenship are as stated below next to my name. In the original, first and sole inventor (if only one nama is listad balow) or and and joint inventor (if plural names are listed below) of the subject matter need and for which a patent is sought on the invention entitled:
	TITLE OF INVENTION
	NTHETIC POLYNUCLEOTIDE CODING FOR HUMAN LACTOFERRIN, ECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT"
(a)[	SPECIFICATION IDENTIFICATION  tion of which: (complete (a), (b) or (c))  ] is attached hereto.  ] was filed on as [ ] U.S. Serial No. 09 /  or [ ] Express Mail No., as Serial No. not yet known, and was amended on (if applicable).

NOTE: Amandments filed after the original papers are deposited with the PTO which contain new matter are not accorded a filing date by being referred to in the declaration. Accordingly, the amendmants involvad ara those filed with the application papers or, in the case of a supplemental declaration, are those amendments claiming matter not encompassed in the original statement of invention or claims. See 37 CFR 1.67.

(c)	[ X ]	was described and claimed in
		PCT International Application No. PCT/IT99/00226
		filed on 19 July 1999 as amended under PCT Article 19 (1)
		on (if anv).

### ACKNOWLEDGMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code Federal Regulations § 1.56.

[ ] In compliance with this duty there is attached an information disclosure statement 37 CFR 1.97.

### PRIORITY CLAIM

I hereby claim foreign priority benefits under Title 35, United States Code, § 119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign applications(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed.

(complete (d) or (e))

- (d) [ ] no such applications have been filed.
- (e) [X] such applications have been filed as follows.

NOTE: Where item (c) is entered above and the International Application which designated the U.S. claimed priority check item (e), enter the details below and make the priority claim.

## EARLIEST FOREIGN APPLICATION(S), IF ANY, FILED WITHIN 12 MONTHS (6 MONTHS FOR DESIGN(S)) PRIOR TO THIS U.S. APPLICATION

COUNTRY	APPLICATION NUMBER	DATE OF FILING (day, month, year)	PRIORITY CLAIMED UNDER 37 USC 119
ITALY	RM98A000478	17 July 1998	[X]YES [ ]NO
			[ ]YES [ ]NO
			[ ]YES [ ]NO
			[ ]YES [ ]NO
			[ ]YES [ ]NO

ALL FOREIGN APPLICATION(S), IF ANY FILEO MORE THAN 12 MONTHS (6 MONTHS FOR OESIGN(S)) PRIOR TO THIS U.S. APPLICATION

#### **POWER OF ATTORNEY**

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

Richard P. Berg, Reg. No. 28,145 Mavis S. Gallenson, Reg. No. 32,464 Kam C. Loule, Reg. No. 33,008 Ross A. Schmitt, Reg. No. 42,529 Victor Repkin, Reg. No. 45,039 John Palmer, Reg. No. 36,885 Peter D. Galloway, Reg. No. 27, 885 William R. Evans, Reg. No. 25, 858

(check the following item, if applicable)

[ ] Attached as part of this declaration and power of attorney is the authorization of the above-named attorney(s) to accept and follow instructions from my representative(s).

#### SEND CORRESPONDENCE TO:

DIRECT TELEPHONE CALLS TO: (Name and telephone number)

Richard P. Berg, Esq. c/o LADAS & PARRY 5670 Wilshire Boulevard, Suite 2100 Los Angeles, California 90036-5679

Richard P. Berg

(323) 934-2300

#### **DECLARATION**

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or Imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

### SIGNATURE(S)

Full name	of sole or first inventor Corrado FOGHER
inventor's	signature
Date	Country of CitizenshipITALY
Residence	Via Ticino, 32, I-26041 Casalmaggiore, ITALY
Post Office	Address(same as residence)
Full name	of second joint inventor, if any
Inventor's	signature
Date	Country of Citizenship
Residence	
Post Office	e Address

# CHECK PROPER BOX(ES) FOR ANY OF THE FOLLOWING ADDED PAGES(S) WHICH FORM A PART OF THIS DECLARATION

Signature for third and subsequent joint inventors. Number of pages added	
[ ] Signature by administrator(trix), executor(trix) or legal representative for deceased incapacitated inventor. Number of pages added	or
[ ] Signature for inventor who refuses to sign or cannot be reached by person authorized under 37 CFR 1.47. Number of pages added Added pages to combined declaration ar power of attorney for divisional, continuation-in-part (CIP) application.  Number of pages added	nd
* * *	
[ ] Authorization of attorney(s) to accept and follow instructions from representative.	
* * *	
If no further pages form a part of this Declaration then end this Declaration with this page and check the following item.	th
[ x ]This declaration ends with this page.	

PATENT

Attorney's	Docket	No. <u>B-40</u>	75PCT 6184 <u>84-4</u>							
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		d :4a.	. I harabu daalara	that						
As a belov	w na <i>m</i> e	ınventoi	r, I hereby declare	: mat.						
				OF DEC			holow)			
	aration is X]	s or the ro original	llowing type: (che	ick one ap	upiicau	ne nem	below)			
-	]	design								
NOTE: If	the dee	supple <i>m</i>	ental s for an Intematior	nal Annlic	ation b	eina file	ed as a d	fivisiona	d. contini	uation or
continuati	ine uec ion-in-pa	art applica	ation, do <u>not</u> checi	k next iter	m; che	ck appr	opriate	one of k	ast three	items.
ſ	Ϋ́	national	stage of P <b>CT</b> ing 3 items apply,							
NOTE: IT	one of t IAL. CO	ne rollow NTINUAT	ing 3 items apply, FION, OR CIP.	uien con	npiete (	anu asc	allacii	MUULU	, , , CEC	,, ,,,,
		divisiona	ıl							
, [ ]		continua	tion tion-in-part (CIP)							
L			INVENTO	Deulo II	DENTI	CIC ATIC	าเก			
WARNIN	G.	If the inv	rentors are each n	of the inv	entors/	of all th	ie claim:	s an exp	danation	of the facts
***	0.	including	g the ownership of	f all the cl	laims a	t the tin	ne the la	st claim	ed inven	ntion was
		made, s	hould be submitte	ed.						
the origin	al, first a names a	and sole i ere listed :	address and citize nventor (if only or below) of the subj	oe name i	is listed	i belowi	or an o	riginai, i	irst and i	ioni mvento
211			TIT	LE OF IN	VENT	ION				
A SYNTH	HETIC P	OLYNU(	CLEOTIDE CODIN	NG FOR	HUMAI	N LACT	OFERR	IN, VEC	CTORS,	CELLS ANI
			SPECIFIC	CATIONI	IDENTI	IFICATI	ON			
the speci	ification	of which:	(complete (a), (b)	) or (c))						
	(a)	[ ]	is attached here	eto.	1 Sori	al No				
	(b)	[]	was filed on or [ ] Express	Mail No.,	, as Se	riel No.	not yet i	known,		
			and was a <i>m</i> end	ed on		(if a	applicab	le).		
	matter a mendi supplei	are not ac ments inv mental de	d after the origina ccorded a filing da olved are those fi claration, are thos at of invention or o	ite by beil led with th se amend	ng rete he app Iments	rrea to i lication ; claiminț	n tne ae papers ( g matter	ciaratio or, in the	an. Accor	raingry, are fa
	(c)	[X]	was described a	and claime	ed in P	'CT Inte	rnationa	I Applic	ation No.	
PCT/IT9	9/00226		filed on 19 July	1000 as a	amenda	ed unde	er PCT 4	vrticle 19	9 (1)	
			on (if any	y) and ide	entified	as U.S.	Serial I	vo. 09/7	43,823	
			<del>-</del>							

### ACKNOWLEDGMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code Federal Regulations § 1.56.

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PRIORITY CLAIM

I hereby claim foreign priority benefits under Title 35, United States Code, § 119 of any foreign application(s) for patent or inventor's certificate or of any PCT international application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign applications(s) for patent or inventor's certificate or any PCT international application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed.

(complete (d) or (e))

(d) [ ] no such applications have been filed.(e) [X] such applications have been filed as follows.

NOTE: Where item (c) is entered above and the International Application which designated the U.S. claimed priority check item (e), enter the details below and make the priority claim.

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COUNTRY	APPLICATION NUMBER	DATE OF FILING (day, month, year)	PRIORITY CLAIMED UNDER 37 USC 119
ITALY	RM98A000478	17 July 1998	[X] YES [] NO
			[]YES []NO
			[]YES []NO
			[] YES [] NO
			[]YES []NO

ALL FOREIGN APPLICATION(S), IF ANY FILED MORE THAN 12 MONTHS (6 MONTHS FOR DESIGN(S)) PRIOR TO THIS U.S. APPLICATION

# CHECK PROPER BOX(ES) FOR ANY OF THE FOLLOWING ADDED PAGES(S) WHICH FORM A PART OF THIS DECLARATION

[ ]	Signature for third and subsequent joint inventors. Number of pages added
[ ] ncapacitated inv	Signature by administrator(trix), executor(trix) or legal representative for deceased or entor. <i>Number of pages added</i>
[ ] under 37 CFR 1./ attorney for divisi	Signature for inventor who refuses to sign or cannot be reached by person authorized 47. Number of pages added Added pages to combined declaration and power of onal, continuation-in-part (CIP) application.  Number of pages added
	***
[ ]	Authorization of attorney(s) to accept and follow instructions from representative.
	***

If no further pages form a part of this Declaration then end this Declaration with this page and check the following item.

[X] This declaration ends with this page.

### POWER OF ATTORNEY

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith. (List name and registration number)

Richard P. Berg, Reg. No. 28,145 Mavis S. Gallenson, Reg. No. 32,464 Victor Repkin, Reg. No. 45,039 John Palmer, Reg. No. 36,885 Peter D. Galloway, Reg. No. 27, 885

Kam C. Louie, Reg. No. 33,008.

William C. Boling, Reg. No. 41,625 William R. Evans, Reg. No. 25, 858 Ross A. Schmitt, Reg. No. 42,529

(check the following item, if applicable)

Attached as part of this declaration and power of attorney is the authorization of the above-[ ] named attorney(s) to accept and follow instructions from my representative(s).

SEND CORRESPONDENCE TO:

DIRECT TELEPHONE CALLS TO: (Name and telephone number)

Richard P. Berg, Esq. c/o LADAS & PARRY 5670 Wilshire Boulevard, Suite 2100 Los Angeles, California 90036-5679

Richard P. Berg

(323) 934-2300

#### DECLARATION

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

### SIGNATURE(S)

Full manner of a sign and fined	inventor Corredo EOGUER	
Full name of sole of first	inventor Corrado FOGHER	
Inventor's signature	Country of Citizenship ITALY	
Date Z. nauce 200	Country of Citizenship HALT	
Residence Casalmaggior	e CR, ITALY	
Post Office Address Via 7	Ticino 32 26041 Casalmaggiore CR ITALY	
Full name of second join	nt inventor, if any	
Inventor's signature		
Date	Country of Citizenship	
		<u> </u>
Residence		
Post Office Address		
Full name of third joint i	nventor, if any	
Inventor's signature		
Date	Country of Citizenship	
Residence		
Post Office Address		

Declaration and Power of Attorney - Page 3 of 4

JC07 Rec'd PCT/PTO 1 6 JAN 20 SEQUENCE LISTING

### GENERAL INFORMATION:

- (i) APPLICANT: PLANTECHNO S.R.L.
- (ii) TITLE OF INVENTION:

SYNTHETIC POLYNUCLEOTIDE ENCODING HUMAN LACTOFERRIN, VECTORS, CELLS AND TRANSGENIC PLANTS CONTAINING IT

- (iii) NUMBER OF SEQUENCES: 22
- (iv) CORRESPONDENCE ADDRESS:
  - (A) ADDRESSEE: Società Italiana Brevetti
  - (B) STREET: Piazza di Pietra, 39
  - (C) CITY: Roma
  - (D) COUNTRY: Italy
  - (E) POSTAL CODE: I-00186
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE:
  - (B) COMPUTER: IBM PC compatible
  - (C) OPERATING SYSTEM: PC-DOS/MS-DOS Rev. 5.0
  - (D) SOFTWARE: Microsoft Word 6.0
- (viii) ATTORNEY INFORMATION
  - (A) NAME: LEONE, Mario (Eng.)
  - (B) REFERENCE: PC-EBR
- (ix) TELECOMMUNICATION INFORMATION
  - (A) TELEPHONE: 06/695441
  - (B) TELEFAX: 06/69544830
  - (C) TELEX: 612287 ROPAT

- (1) INFORMATION FOR SEQ ID NO: 1:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 2079 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (ix) FEATURES
  - (A) NAME: LFUSYN
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GGC CGT AGG AGA AGG AGT GTT CAA TGG TGC GCA GTA TCA CAA CCA 45 G R R R R S V O W C A V S O P

GAG GCC ACA AAA TGC TTC CAA TGG CAA AGG AAT ATG AGA AAA GTT 90 E A T K C F Q W Q R N M R K V

CGT GGA CCT CCT GTA TCT TGC ATA AAG AGA GAT TCA CCC ATC CAG 135
R G P P V S C I K R D S P I Q

TGT ATC CAG GCA ATT GCG GAA AAC AGA GCT GAT GCT GTG ACT CTT 180

C I Q A I A E N R A D A V T L

GAT GGT GGT TTC ATA TAC GAG GCA GGA CTT GCC CCA TAC AAA CTG 225 D G G F I Y E A G L A -P Y K L

CGA CCT GTA GCG GCG GAA GTC TAC GGG ACC GAA AGA CAA CCA CGA 270
R P V A A E V Y G T E R Q P R

ACT CAC TAT TAT GCT GTG GCT GTT GTG AAG AAG GGC GGA TCT TTT 315
T H Y Y A V A V V K K G G S F

CAG CTG AAC GAA CTT CAA GGT CTG AAG TCA TGC CAC ACA GGA CTT 360
Q L N E L Q G L K S C H T G L

CGC AGG ACC GCT GGA TGG AAT GTC CCT ATA GGG ACA CTT CGT CCA 405

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TTC TTG AAT TGG ACG GGT CCA CCT GAG CCC ATT GAG GCA GCT GTG 450 Τ G Ρ Ρ E P I Е Α Α GCA AGA TTC TCA GCC TCT TGT GTT CCA GGT GCA GAT AAA GGA 495 R F E S Α S C V P G Α D K G CAA TTC CCC AAC CTT TGT CGC CTG TGT GCG GGG ACA GGG GAA AAC 540 R L С Ρ N L С Α G T G Ε Ν AAA TGT GCA TTC TCA TCC CAG GAA CCG TAC TTC AGC TAC TCT GGT 585 С Ė Α F S S Q P Y Ę S Y S G GCC TTT AAG TGT CTT AGA GAC GGT GCT GGA GAT GTT GCT TTT ATT C L R D G Α G D Α FI AGA GAG AGC ACA GTG TTT GAG GAT CTT TCA GAC GAG GCT GAA AGG 675 S T V F Ė D L S DEA Ė GAC GAG TAT GAG TTA CTC TGC CCA GAC AAC ACT CGT AAG CCA GTT Ε Y Ė L L C P D N T R K GAC AAG TTC AAA GAT TGC CAT CTT GCA CGG GTC CCT TCT CAT GCC V P S Н Α K D С Н L A R GTT GTG GCA CGA AGT GTT AAT GGA AAG GAG GAT GCC ATC TGG AAT 810 E D K Ι V A R S V N G Α CTT CTC CGC CAA GCA CAG GAA AAG TTT GGA AAG GAC AAG TCA CCG L R Q Α QE K F G K K P AAA TTC CAG CTC TTT GGT TCC CCT AGT GGG CAG AAA GAT CTT CTG 900 F L F G S P S G Q K D TTC AAG GAC TCT GCC ATT GGG TTT TCG AGA GTG CCA CCT AGG ATA 945

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GAT TCT GGG TTG TAC CTT GGC TCC GGA TAC TTT ACT GCA ATT CAG 990 S G L Ļ G S G Υ F Т Α AAC TTG AGG AAA AGT GAG GAG GAA GTT GCT GCC CGG CGT GCG CGG 1035 E Ē V A Α R R GTC GTT TGG TGT GCG GTG GGA GAG CAA GAG TTG CGC AAG TGT AAC 1080 С Α ٧ G Ε Ε Q CAG TGG AGT GGT TTG AGC GAA GGA TCT GTG ACC TGC TCA TCG GCC 1125 G Ε G S V Т С S TCC ACT ACA GAA GAT TGC ATC GCC CTG GTG TTG AAA GGA GAA GCT 1170 CI Α v E D L L K G E GAT GCC ATG AGT TTG GAT GGA GGA TAT GTT TAC ACT GCA GGT AAA 1215 D Α S L Ð G G Y ٧ Y Т Α G TGT GGT TTG GTG CCT GTC CTT GCA GAG AAC TAC AAA TCA CAA CAA V P V L E N Y K S Q C G L A AGC AGT GAC CCT GAT CCT AAC TGT GTG GAT AGA CCT GTG GAA GGA 1305 С V D V S D P D P N R P E G TAT CTT GCT GTG GCG GTG GTT AGG AGA TCA GAC ACT AGC CTT ACC 1350 Α V Α ٧ V R R S Ď T S L TGG AAC TCT GTG AAA GGC AAG AAG TCC TGC CAC ACC GCC GTG GAC V S v K G K K S С Н Т Α D AGG ACT GCA GGT TGG AAT ATC CCC ATG GGA TTG CTC TTC AAC CAG T Α G W NI P M G L L F N Q ACG GGC TCC TGC AAA TTT GAT GAA TAT TTC AGT CAA AGC TGT GCC 1485 G S С K F E Y F S Q S Ç D CCT GGT TCT GAC CCA AGA TCT AAT CTC TGT GCT TTG TGT ATT GGA

P	G	s	D	P	R	S	N	L	С	A	L	С	I	G	
													GAG E	AGA R	1575
	TAC Y													GCT A	1620
	GAC D													ACT T	1665
	GGA G							GCT A						GCA A	1710
	TTT F												T GTO	G ACT	1755
gaa		AGA	AGC			CTT	GCC		GCC	CCG	AAT	CAT		GTG V	1800
GTG		CGT	ATG	GAT	AAG	GTG	gaa	CGC	TTG	AAA	CAG	GTG		CTC	1845
CAC	CAA	CAG	GCT	AAA	TTT	GGT	AGA	AAT	GGA	TCT	GAC	TGC	CCG	GAC	1890
AAG	TTT	TGC	TTA	TTC	CAG	TCT	GAA	ACC	AAA	AAC	CTT	TTG	P TTC	AAT	1935
GAC	AAC	ACT	GAG	TGT	CTT	GCC	AGA	CTC	CAT	GGC	AAA	ACA		TAT	1980
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AGG AAG TAA R K *		2079
(2) INFORMATION FOR SEQ ID NO: 2:		
(i) SEQUENCE CHARACTERISTICS		
(A) LENGTH: 30 bp		
(B) TYPE: nucleic acid		
(C) STRANDEDNESS: single		
(D) TOPOLOGY: linear		
(ii) MOLECULE TYPE: oligonucleotide		
(ix) FEATURES		
(A) NAME: PLT 46		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:		
GGATCCATGG GCCGTAGGAG AAGGAGTGTT	3	0
(3) INFORMATION FOR SEQ ID NO: 3:		
(i) SEQUENCE CHARACTERISTICS		
(A) LENGTH: 32 bp		
(B) TYPE: nucleic acid		
(C) STRANDEDNESS: single		
(D) TOPOLOGY: linear		
(ii) MOLECULE TYPE: oligonucleotide		
(ix) FEATURES		
(A) NAME: PLT 47		
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:		
GAGCTCCTTC GGTTTTACTT CCTGAGGAAT TC	32	
(4) INFORMATION FOR SEQ ID NO: 4:		
(i) SEQUENCE CHARACTERISTICS		
(A) LENGTH: 42 bp		
(B) TYPE: nucleic acid		

(C) STRANDEDNESS: single

(ii) MOLECULE TYPE: oligonucleotide

(D) TOPOLOGY: linear

(ix) FEATURES

(A)	NAME:	PLT	48
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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

### TCTAGATAAA ATAATCTATA CATTAAAAAA TTTGATTTTA AA 42

- (5) INFORMATION FOR SEQ ID NO: 5:
  - (i) SEQUENCE CHARACTERISTICS
    - (A) LENGTH: 36 bp
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: oligonucleotide
  - (ix) FEATURES
    - (A) NAME: PLT 49
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

### GGATCCGACT GAGTCGGATA AGAAGAAAAG AAAAGA 36

- (6) INFORMATION FOR SEQ ID NO: 6:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 36 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 50
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

### TCTAGAGTTT TCAAATTTGA ATTTTAATGT GTGTTG 36

- (7) INFORMATION FOR SEQ ID NO: 7:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 36 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single

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- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 51
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

GGATCCCACC TTAAGGAGGT TGCAACGAGC GTGGCA 36

- (8) INFORMATION FOR SEQ ID NO:8:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 250 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 120
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

GGC CGT AGG AGA AGG AGT GTT CAA TGG TGC GCA GTA TCA CAA CCA GAG
GCC ACA AAA TGC TTC CAA TGG CAA AGG AAT ATG AGA AAA GTT CGT GGA
CCT CCT GTA TCT TGC ATA AAG AGA GAT TCA CCC ATC CAG TGT ATC CAG
GCA ATT GCG GAA AAC ĀGA GCT GAT GCT GTG ACT CTT GAT GGT GGT TTC
ATA TAC GAG GCA GGA CTT GCC CCA TAC AAA CTG CGA CCT GTA GCG GCG
GAA GTC TAC G

- (9) INFORMATION FOR SEQ ID NO:9:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 250 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 121
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

GC ACC TGG AAC ACA AGA GGC TGA GAA GAA TCT TGC CAC AGC TGC CTC

AAT GGG CTC AGG TGG ACC CGT CCA ATT CAA GAA TGG ACG AAG TGT CCC

TAT AGG GAC ATT CCA TCC AGC GGT CCT GCG AAG TCC TGT GTG GCA TGA

CTT CAG ACC TTG AAG TTC GTT CAG CTG AAA AGA TCC GCC CTT CTT CAC

AAC AGC CAC AGC ATA ATA GTG AGT TCG TGG TTG TCT TTC GGT CCC GTA

GAC TTC CGC CG

- (10) INFORMATION FOR SEQ ID NO:10:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 250 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 122
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

AAC AGA GTA GCT GAA GTA GTC TGG GCA GAG TAA CTC ATA CTC GTC

CCT TTC AGC CTC GTC TGA AAG ATC CTC AAA CAC TGT GCT CTC

AAT AAA AGC AAC ATC TCC AGC ACC GTC TCT AAG ACA CTT AAA GGC

ACC AGA GTA GCT GAA GTA CGG TTC CTG GGA TGA GAA TGC ACA TTT

GTT TTC CCC TGT CCC CGC ACA CAG GCG ACA AAG GTT GGG GAA TTG

TCC TTT ATC TGC ACC TGG AAC ACA A

- (11) INFORMATION FOR SEQ ID NO:11:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 255 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 123
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

GTA CAA CCC AGA ATC TAT CCT AGG TGG CAC TCT CGA AAA CCC AAT GGC

AGA GTC CTT GAA CAG AAG ATC TTT CTG CCC ACT AGG GGA ACC AAA GAG

CTG GAA TTT CGG TGA CTT GTC CTT TCC AAA CTT TTC CTG TGC TTG GCG

GAG AAG ATT CCA GAT GGC ATC CTC CTT TCC ATT AAC ACT TCG TGC CAC

AAC GGC ATG AGA AGG GAC CCG TGC AAG ATG GCA ATC TTT GAA CTT GTC

AAC TGG CTT ACG AGT

- (12) INFORMATION FOR SEO ID NO:12:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 251 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 124
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

- (13) INFORMATION FOR SEQ ID NO:13:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 75 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
- (D) OTHER INFORMATION: leader sequence of protein 7S basic globulin
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:
    ATGGCTTCTA TCCTCCACTA CTTTTTAGCC CTCTCTTT CTTGCTCTTT 50
    TCTTTTCTTC TTATCCGACT CAGTC 75
  - (14) INFORMATION FOR SEQ ID NO:14:
  - (i) SEQUENCE CHARACTERISTICS
    - (A) LENGTH: 191 bp
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: double
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: oligonucleotide
  - (ix) FEATURES
  - (D) OTHER INFORMATION: leader sequence of protein  $\beta\text{-}$  conglycinine
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

ATGATGAGAG CGCGGTTCCC ATTACTGTTG CTGGGAGTTG TTTTCCTAGC 50
ATCAGTTTCT GTCTCATTTG GCATTGCGTA TTGGGAAAAG CAGAACCCCA 100
GTCACAACAA GTGCCTCCGA AGTTGCAATA GCGAGAAAGA CTCCTACAGG 150
AACCAAGCAT GCCACGCTCG TTGCAACCTC CTTAAGGTG 189

- (15) INFORMATION FOR SEQ ID NO:15:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 250 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 125
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

  GAGCAATCCC ATGGGGATAT TCCAACCTGC AGTCCTGTCC ACGGCGGTGT 50

  GGCAGGACTT CTTGCCTTTC ACAGAGTTCC AGGTAAGGCT AGTGTCTGAT 100

  CTCCTAACCA CCGCCACAGC AAGATATCCT TCCACAGGTC TATCCACACA 150

  GTTAGGATCA GGGTCACTGC TTTGTTGTGA TTTGTAGTTC TCTGCAAGAC 200

  AGGCACCAAA CCACATTTAC CTGCAGTGTA AACATATCCT CCATCCAAAC 250
  - (16) INFORMATION FOR SEQ ID NO:16:
  - (i) SEQUENCE CHARACTERISTICS
    - (A) LENGTH: 254 bp
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: oligonucleotide
  - (ix) FEATURES
    - (A) NAME: PLT 126
- (xì) SEQUENCE DESCRIPTION: SEQ ID NO:16:

  CCATCAGTGT TCTGCAAGAC AGTGACATCT TTCACAAATG CAACGTCTCC 50

  AGCATTCTCA GCCAAGCAAC GGAAAGCCCC AGTGTAACCG TAGTATCTCT 100

  CGTTGCTGTT GGGAACGCAC TTATTCTCAC CTTGCTCATC TCCAATACAC 150

  AAAGCACAGA GATTAGATCT TGGGTCAGAA CCAGGGGCAC AGCTTTGACT 200

  GAAATATTCA TCAAATTTGC AGGAGCCCGT CTGGTTGAAG AGCAAGCCCA 250

  TGGG

- (17) INFORMATION FOR SEQ ID NO:17:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 229 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
  - (ix) FEATURES
    - (A) NAME: PLT 127
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:
    GCAGTCAGAT CCATTTCTAC CAAATTTAGC CTGTTGGTGG AGCAACACCT 50
    GTTTCAAGCG TTCCACCTTA TCCATACGAG ACACCACAGC ATGATTCGGG 100
    GCCATGGCAA GATGGCAGCT TCTAGCTTCA GTCACAGGCT TACGTTTGCC 150
    ATCGAGGCAC AGCAACGCAA AGTCTGCAAG CTTCAAATCC TTAGCCCATG 200
    CCTCATTGTT ATTTCCATCA GTGTTCTGC 229
  - (18) INFORMATION FOR SEQ ID NO:18:
  - (i) SEQUENCE CHARACTERISTICS
    - (A) LENGTH: 210 bp
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: oligonucleotide
  - (ix) FEATURES
    - (A) NAME: PLT 128
  - (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

    TTACTTCCTT AGGAATTCAC AGGCTTCTAG GAGTGGGGAG GTTGAGCACT 50

    TTTTCAGATT AGTAATGCCT GCGACATACT GTGGTCCCAA ATATTTTCA 100

    TATGTTGTTT TGCCATGGAG TCTGGCAAGA CACTCAGTGT TGTCATTGAA 150

    CAAAAGGTTT TTGGTTTCAG ACTGGAATAA GCAAAACTTG TCCGGGCAGT 200

    CAGATCCATT 210
  - (19) INFORMATION FOR SEQ ID NO:19:
  - (i) SEQUENCE CHARACTERISTICS
    - (A) LENGTH: 30-bp
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: oligonucleotide

{	ix	)	FEAT	ľ	ΙR	E.S

- (A) NAME: PLT 129
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

### GGATCCATGG GCCGTAGGAG AAGGAGTGTT

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- (20) INFORMATION FOR SEQ ID NO: 20:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: 28 bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: oligonucleotide
- (ix) FEATURES
  - (A) NAME: PLT 120
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

GAGCTCTTAC TTCCTTAGGA ATTCACAG

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- (21) INFORMATION FOR SEQ ID NO:21:
- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (ix) FEATURES
- (D) OTHER INFORMATION: sequence of promoter of gene encoding 7S basic globulin
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

TAAAATAATC TATACATTAA AAAATTTGAT TTTAAAATTT TAGAAATTCA 50

TGATTTTATT TTTTTTTACC AGAAATCCGT TAATATTGTT AAAATATTAC 100

CAACTAATTT ATAAATTTTA TTTTAAGGCA ATTAAGCATG TTTGATAAAA 150

TATATATAT GTTATAAATA CTTTTCAAAA GTATAAAGTT GATGATGGCG 200

TGGTGGTAGA TTATTTTAGT TCTAGGTTCG AATGCAAGTT GGTTTAGACA 250

TTTAGCCTTA TTCTTTTTC TAACCAAAAT AAATGTAAAT GGAAAACCTT 300

TAGGAAAAA AAGAAATCAA AATTGAAAAC ATCATCCGGT GGAGTCGAGA 350

AGCCCACACC CACGTGACCC AACAATATTA AAATAAGAGT TTGCTCTACA 400

GTAAATGCGA	TACTTTTTTA	TTCAATACTT	TTTCCACTTC	TAAAATCTTG	450
GAGATTTGCA	CCGTTAACTA	ATTAAGTGTT	ATATCCAACG	GTCCTAAAAA	500
AACTTGTGTA	CCGTGCCTCA	CATTTCAACT	TTGCGCACCC	TGAAAGCCGT	550
TATGTTTAGG	TTAGTGTTTG	CAACAGTTGA	AGCGCATCAC	TCAGGAGGCT	600
ACTTGGTCTT	GCTTTTGCGT	CTTTTGTTCA	ATTTTTCACG	TGATTTTGTT	650
GGTGAACACG	CGTACTTGAA	ACTTATTATA	AATTACATAA	TTTTATAAGT	700
TTCACTTCTT	ATATAATACT	CATATAATAT	ATAGGGTTTA	GAATGCCAAT	750
TTTTAAAAAA	AGAATAAAAA	AATAAATAGA	ATAAAATCGA	AAAAATGAAA	800
TGTAAAAAAT	TTGAGGGGGA	CAAATAAAAT	ATGAAAGTCT	ATTATTTAAA	850
TTTTCCATTA	GAATTCTATT	TTCCTTAGTT	AATATGAGCT	AGCCAGTTGG	900
GAGATACACG	AAAATGTCAT	GAAACAGTTG	CATGTAGGGA	AATTAATGTA	950
GTAGAGGGAT	AGCAAGACAA	AAATCCAAGC	CAAGCTAGCT	GCTCACGCGA	1000
ACTCGATCCA	CACGTCCTTT	ACAGAGTTTC	AAACGGATGA	AATCTGCATG	1050
GCATGCAACT	AAAGCATTGT	TCTCAGCTGC	CAAGTACCCC	TCACACTCAC	1100
CAACCCTTTG	TTTTTCTCCC	CATTGCATGT	TAACTCAAGT	TTATCCTTTC	1150
TTTGCTTCTG	GAAATTTCAC	AAGCCTCAAA	CACGTCGACG	TCCAATCTTG	.1200
TGACCAACAC	GGCCAAAAGA	AAAGAGAATC	TCATCCCGTT	CACACTTAGC	1250
CACTTAAAGC	TAGCCAAACG	GTGATCTTTC	TCTATATATT	GTAGCTCTCT	1300
AACACAACCA	ACACTACCAT	TATTCAATAT	TCAAACCTTG	CTCTATACTA	1350
CACACACTAG	AAGAATA				1367

### (22) INFORMATION FOR SEQ ID NO:22:

- (i) SEQUENCE CHARACTERISTICS
  - (A) LENGTH: bp
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: double
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (ix) FEATURES
- (D) OTHER INFORMATION: sequence of promoter of gene coding for  $\beta\text{--}\text{conglycinine}$
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:
- GTTTTCAAAT TTGAATTTTA ATGTGTGTTG TAAGTATAAA TTTAAAATAA 50
- AAATAAAAC AATTATTATA TCAAAATGGC AAAAACATTT AATACGTATT 100
- ATTTATTAAA AAAATATGTA ATAATATATT TATATTTTAA TATCTATTCT 150
- TATGTATTT TTAAAAATCT ATTATATATT GATCAACTAA AATATTTTTA 200
  - TATCTACACT TATTTTGCAT TTTTATCAAT TTTCTTGCGT TTTTTGGCAT 250

TAATAATTTA	GACTATTCTT	TAATAATCAA	TCATTATTCT	TACATGGTAC	300
ATATTGTTGG	AACCATATGA	AGTGTTCATT	GCATTTGACT	ATGTGGATAG	350
TGTTTTGATC	CATGCCCTTC	ATTTGCCGCT	ATTAATTAAT	TTGGTAACAG	400
ATTCGTTCTA	ATCAGTTACT	TAATCCTTCC	TCATCATAAT	TAATCTGGTA	450
GTTCGAATGC	CATAATATTG	ATTAGTTTTT	TGGACCATAA	GAAAAAGCCA	500
AGGAACAAAA	GAAGACAAAA	CACAATGAGA	GTATCCTTTG	CATAGCAATG	550
TCTAAGTTCA	TAAAATTCAA	ACAAAAACGC	AATCACACAC	AGTGGACATC	600
ACTTATCCAC	TAGCTGATCA	GGATCGCCGC	GTCAAGAAAA	AAAAACTGGA	650
CCCCAAAAGC	CATGCACAAC	AACACGTACT	CACAAAGGCG	TCAATCGAGC	700
AGCCCAAAAC	ATTCACCAAC	TCAACCCATC	ATGAGCCCAC	ACATTTGTTG	750
TTTCTAACCC	AACCTCAAAC	TCGTATTCTC	TTCCGCCACC	TCATTTTTGT	800
TTATTTCAAC	ACCCGTCAAA	CTGCATCCCA	CCCCGTGGCC	AAATGTTCAT	850
GCATGTTAAC	AAGACCTATG	ACTATAAATA	TCTGCAATCT	CGGCCCAAGT	900
TTTCATCATC	AAGAACCAGT	TCAATATCCT	AGTACGCCGT	ATTAAAGAAT	950
TTAAGATATA	CT				962

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